PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5: C12N 15/00, C07K 7/10, 13/00 A61K 37/02, 27/00

A1

(11) International Publication Number:

WO 92/07073

(43) International Publication Date:

30 April 1992 (30.04.92)

(21) International Application Number:

PCT/US91/07635

(22) International Filing Date:

18 October 1991 (18.10.91)

(30) Priority data:

599,543

18 October 1990 (18.10.90) US

(71) Applicant: CREATIVE BIOMOLECULES, INC. [US/US]; 35 South Street, Hopkinton, MA 01748 (US).

(72) Inventors: OPPERMANN, Hermann; 25 Summer Hill Road, Medway, MA 02053 (US). OZKAYNAK, Engin; 44 Purdue Drive, Milford, MA 01757 (US). RUEGER, David, C.; 150 Edgemere Road, Apt. 4, West Roxbury, MA 02132 (US). KUBERASAMPATH, Thangavel; 6 Spring Street, Medway, MA 02053 (US).

(74) Agent: PITCHER, Edmund, R.; Testa, Hurwitz & Thibeault, Exchange Place, 53 State Street, Boston, MA 02109-2809 (US).

(81) Designated States: AT (European patent), AU, BE (European patent), CA, CH (European patent), DE (European patent), DK (European patent), ES (European patent), FR (European patent), GB (European patent), GR (European patent), IT (European patent), JP, LU (European patent), NL (European patent), SE (European patent).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: OSTEOGENIC PEPTIDES

(57) Abstract

Disclosed are 1) the cDNA and amino acid sequences for novel polypeptide chains useful as subunits of dimeric osteogenic proteins, 2) osteogenic devices comprising these proteins in association with an appropriate carrier matrix, 3) methods of producing the polypeptide chains using recombinant DNA technology, and 4) methods of using the osteogenic devices to mimic the natural course of endochondral bone formation in mammals.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	ES	Spain	MG	Madagascar
AU	Australia	FI	Finland	ML	Mali
BB	Barbados	FR	France	MN	Mongolia
BE	Belgium	GA	Gabon	MR	Mauritania
BF	Burkina Faso	GB	United Kingdom	MW	Malawi
BG	Bulgaria	GN	Guinca	NL	Netherlands
BJ	Benin	GR	Greece	NO	Norway
BR	Brazil	HU	Hungary	PL	Poland
CA	Canada	IT	Italy	RO	Romania
CF	Central African Republic	JP	Japan	SD	Sudan
CG	Congo	KP	Democratic People's Republic	SE	Sweden
СН	Switzerland		of Korea	SN	Senegal
Cl	Côte d'Ivoire	KR	Republic of Korea	su+	Soviet Union
CM	Cameroon	LI	Liechtenstein	TD	Chad
cs	Czechoslovakia	LK	Sri Lanka	TG	Togo
DE.	Germany	LU	Luxembourg	US	United States of America
DK	Denmark	MC	Monaco		

⁺ Any designation of "SU" has effect in the Russian Federation. It is not yet known whether any such designation has effect in other States of the former Soviet Union.

Osteogenic peptides

Background of the Invention

This invention relates to novel polypeptide chains and to osteogenic proteins comprising these polypeptide chains which are capable of inducing osteogenesis in mammals; to genes encoding the polypeptide chains; to methods for their production using recombinant DNA techniques, and to bone and cartilage repair procedures using the osteogenic proteins.

Mammalian bone tissue is known to contain one or more proteinaceous materials, presumably active during growth and natural bone healing, which can induce a developmental cascade of cellular events resulting in endochondral bone formation. This active factor (or. factors) has variously been referred to in the literature as bone morphogenetic or morphogenic protein, bone inductive protein, osteogenic protein, osteogenin, or osteoinductive protein.

The developmental cascade of bone differentiation consists of recruitment of mesenchymal cells,

25 proliferation of progenitor cells, calcification of cartilage, vascular invasion, bone formation, remodeling, and finally marrow differentiation (Reddi (1981) Collagen Rel. Res. 1:209-226).

Though the precise mechanisms underlying these phenotypic transformations are unclear, it has been shown that the natural endochondral bone differentiation activity of bone matrix can be dissociatively extracted and reconstituted with

05

10

15

20

30

inactive residual collagenous matrix to restore full bone induction activity (Sampath and Reddi, (1981)

Proc. Natl. Acad. Sci. USA 78:7599-7603). This provides an experimental method for assaying protein extracts for their ability to induce endochondral bone in vivo. Several species of mammals produce closely related protein as demonstrated by cross species implant experiments (Sampath and Reddi (1983) Proc. Natl. Acad. Sci. USA 80:6591-6595).

The potential utility of these proteins has been recognized widely. It is contemplated that the availability of the protein would revolutionize orthopedic medicine, certain types of plastic surgery, and various periodontal and craniofacial reconstructive procedures.

The observed properties of these protein fractions have induced an intense research effort in various laboratories directed to isolating and identifying the pure factor or factors responsible for osteogenic activity. The current state of the art of purification of osteogenic protein from mammalian bone is disclosed by Sampath et al. ((1987) Proc. Natl. Acad. Sci. USA 84: 7109-7113). Urist et al. (1984) Proc. Soc. Exp. Biol. Med. 173: 194-199 disclose a human osteogenic protein fraction which was extracted from demineralized cortical bone by means of a calcium chloride-urea inorganic-organic solvent mixture, and retrieved by differential precipitation in quanidine-hydrochloride and preparative gel electrophoresis. The authors report that the protein fraction has an amino acid composition of an acidic polypeptide and a molecular weight in a range of 17-18 kD.

Urist et al. (1984) Proc. Natl. Acad. Sci. USA 81: 371-375 disclose a bovine bone morphogenetic protein extract having the properties of an acidic polypeptide and a molecular weight of approximately 18 kD. The of authors reported that the protein was present in a fraction separated by hydroxyapatite chromatography, and that it induced bone formation in mouse hindquarter muscle and bone regeneration in trephine defects in rat and dog skulls. Their method of obtaining the extract 10 from bone results in ill-defined and impure preparations.

European Patent Application Serial No. 148,155, published October 7, 1985, purports to disclose osteogenic proteins derived from bovine, porcine, and 15 human origin. One of the proteins, designated by the inventors as a P3 protein having a molecular weight of 22-24 kD, is said to have been purified to an essentially homogeneous state. This material is reported to induce bone formation when implanted into 20 animals.

International Application No. PCT/087/01537, published January 14, 1988, discloses an impure fraction from bovine bone which has bone induction qualities. The named applicants also disclose putative 25 "bone inductive factors" produced by recombinant DNA techniques. Four DNA sequences were retrieved from human or bovine genomic or cDNA libraries and expressed in recombinant host cells. While the applicants stated that the expressed proteins may be bone morphogenic proteins, bone induction was not demonstrated, suggesting that the recombinant proteins are not osteogenic. The same group reported subsequently (Science, 242:1528, Dec, 1988) that three of the four

factors induce cartilage formation, and postulate that bone formation activity "is due to a mixture of regulatory molecules" and that "bone formation is most likely controlled ... by the interaction of these of molecules." Again, no bone induction was attributed to the products of expression of the cDNAs. See also Urist et al., EPO,212,474 entitled Bone Morphogenic Agents.

Wang et al. (1988) Proc. Nat. Acad. Sci. USA 85:

10 9484-9488 discloses the purification of a bovine bone morphogenetic protein from guanidine extracts of demineralized bone having cartilage and bone formation activity as a basic protein corresponding to a molecular weight of 30 kD determined from gel elution.

15 Purification of the protein yielded proteins of 30, 18 and 16 kD which, upon separation, were inactive. In view of this result, the authors acknowledged that the exact identity of the active material had not been determined.

Wang et al. (1990) <u>Proc. Nat. Acad. Sci. USA</u> 87: 2220-2227 describes the expression and partial purification of one of the cDNA sequences described in PCT 87/01537. Consistent cartilage and/or bone formation with their protein requires a minimum of 600 25 ng of 50% pure material.

International Application No. PCT/89/04458

published April 19, 1990 (Int. Pub. No. WO90/003733),

describes the purification and analysis of a family of
 osteogenic factors called "P3 OF 31-34". The protein

30 family contains at least four proteins, which are
 characterized by peptide fragment sequences. The
 impure mixture P3 OF 31-34 is assayed for osteogenic

- 5 -

 05 activity. The activity of the individual proteins is neither assessed nor discussed.

It is an object of this invention to provide novel polypeptide chains useful as subunits of dimeric osteogenic proteins capable of endochondral bone formation in allogenic and xenogenic implants in mammals, including humans. Another object is to provide genes encoding these polypeptide chains and methods for the production of osteogenic proteins comprising these polypeptide chains using recombinant DNA techniques, as well as to provide antibodies capable of binding specifically to these proteins.

These and other objects and features of the invention will be apparent from the description, drawings, and claims which follow.

Summary of the Invention

This invention provides novel polypeptide chains useful as either one or both subunits of dimeric osteogenic proteins which, when implanted in a 05 mammalian body in association with a matrix, can induce at the locus of the implant the full developmental cascade of endochondral bone formation and bone marrow differentiation.

A key to these developments was the elucidation of amino acid sequence and structure data of native bovine osteogenic protein. A protocol was developed which results in retrieval of active, substantially pure osteogenic protein from bovine bone having a half-maximum bone forming activity of about 0.8 to 1.0 ng per mg of implant. The availability of the material enabled the inventors to elucidate key structural details of the protein necessary to achieve bone formation. Knowledge of the protein's amino acid sequence and other structural features enabled the identification and cloning of native genes in the human genome.

Consensus DNA sequences based on partial sequence data and observed homologies with regulatory proteins disclosed in the literature were used as probes for extracting genes encoding osteogenic protein from human genomic and cDNA libraries. One of the consensus sequences was used to isolate a previously unidentified gene which, when expressed, encoded a protein comprising a region capable of inducing endochondral bone formation when properly modified, incorporated in a suitable matrix, and implanted as disclosed herein. The gene, called "hOP1" or "OP-1" (human OP-1), is

described in greater detail in copending U.S. 422,699, the disclosure of which is herein incorporated by reference.

In its native form, hOP1 expression yields an 05 immature translation product ("hOP1-PP", where "PP" refers to "prepro form") of about 400 amino acids that subsequently is processed to yield a mature sequence of 139 amino acids ("OP1-18"). The active region (functional domain) of the protein comprises the 10 C-terminal 97 amino acids of the hOP1 sequence ("OPS"). A long active sequence is OP7 (comprising the C-terminal 102 amino acids).

Further probing of mammalian cDNA libraries (human and mouse) with sequences specific to hOP1 also has 15 identified novel OP1-like sequences herein referred to as "OP2" ("hOP2" or "mOP2"). The OP2 proteins share significant amino acid sequence homology, approximately 74%, with the active region of the OP1 proteins (e.g., OP7), and less homology with the intact mature form 20 (e.g., OP1-18, 58% amino acid homology).

The amino acid sequence of the osteogenic proteins disclosed herein also share significant homology with various of the regulatory proteins on which the consensus probe was modeled. In particular, the 25 proteins share significant homology in their C-terminal sequences, which comprise the active region of the osteogenic proteins. (Compare, for example, OP7 with DPP from Drosophila and Vgl from Xenopus. See, for example, U.S. Pat. No. 5,011,691). In addition, these 30 proteins share a conserved six or seven cysteine skeleton in this region (e.g., the linear arrangement of these C-terminal cysteine residues is conserved in

the different proteins.) See, for example, OP7, whose sequence defines the seven cysteine skeleton, or OPS, whose sequence defines the six cysteine skeleton. The OP2 proteins also contain an additional cysteine 05 residue within this region.

Thus, in one preferred aspect, the invention comprises osteogenic proteins comprising a polypeptide chain comprising an amino acid sequence described by Seq. ID No. 3 or 5, including allelic and species variants thereof, and naturally-occurring or biosynthetic mutants, such that a dimeric protein comprising this polypeptide chain has a conformation capable of inducing endochondral bone formation when implanted in a mammal in association with a suitable matrix. Useful proteins include the full-length protein, mature proteins and truncated proteins comprising the functional domain described by the C-terminal.

In addition, the invention is not limited to thse

20 specific constructs. Thus, the osteogenic proteins of
this invention comprising any of these polypeptide
chains may include forms having varying glycosylation
patterns, varying N-termini, a family of related
proteins having regions of amino acid sequence homology
which may be naturally occurring or biosynthetically
derived, and active truncated or mutated forms of the
native amino acid sequence, produced by expression of
recombinant DNA in procaryotic or eucaryotic host
cells. Active squances useful as osteogenic proteins
of this invention are envisioned to include proteins
capable of inducing endochondral bone formation when
implanted in a mammal in association wiht a matrix and
having at lest a 70% sequence homology, preferably at

- 9 -

least 80%, with the amino acid sequence of OPS. This includes longer forms of a given protein, as well as allelic variants and muteins, including addition and deletion mutants, such as those which may alter the conserved C-terminal cysteine skeleton, provided that the alteration still allows the protein to form a dimeric species having a conformation capable of inducing bone formation in a mammal when implanted in the mammal in association with a matrix.

The novel polypeptide chains and the osteogenic roteins they comprise can be expressed from intact or truncated cDNA or from synthetic DNAs in procaryotic or eucaryotic host cells, and then purified, cleaved, refolded, dimerized, and implanted in experimental animals. Currently preferred host cells include E.coli or mammalian cells, such as CHO, COS or BSC cells. The osteogenic protein of the invention may include forms having varying glycosylation patterns, varying N-termini, a family of related proteins having regions of amino acid sequence homology, and active truncated or mutated forms of native or biosynthetic proteins, produced by expression of recombinant DNA in host cells.

Thus, in view of this disclosure, skilled genetic
25 engineers can isolate genes from cDNA or genomic
libraries of various different species which encode
appropriate amino acid sequences, or construct DNAs
from oligonucleotides, and then can express them in
various types of host cells, including both procaryotes
30 and eucaryotes, to produce large quantities of active
proteins capable of inducing bone formation in mammals
including humans. In view of this disclosure, those
skilled in the art, using standard immunology

techniques also may create antibodies capable of binding specifically to the osteogenic proteins disclosed herein, including fragments thereof.

The osteogenic proteins are useful in clinical applications in conjunction with a suitable delivery or 05 support system (matrix). The matrix is made up of particles of porous materials. The pores must be of a dimension to permit progenitor cell migration and subsequent differentiation and proliferation. particle size should be within the range of 70 - 850 mm, preferably 150mm - 420mm. It may be fabricated by close packing particulate material into a shape spanning the bone defect, or by otherwise structuring as desired a material that is biocompatible (noninflammatory) and, biodegradable in vivo to serve as a 15 "temporary scaffold" and substratum for recruitment of migratory progenitor cells, and as a base for their subsequent anchoring and proliferation. Currently preferred carriers include particulate, demineralized, 20 guanidine extracted, species-specific (allogenic) bone, and specially treated particulate, protein extracted, demineralized, xenogenic bone. Optionally, such xenogenic bone powder matrices also may be treated with proteases such as trypsin and/or fibril modifying 25 agents to increase the intraparticle intrusion volume and surface area. Useful agents include solvents such as dichloromethane, trichloroacetic acid, acetonitrile and acids such as trifluoroacetic acid and hydrogen fluoride. Alternatively, the matrix may be treated $_{
m 30}$ with a hot aqueous medium having a temperature within the range of about 37°C to 75°C, including a heated acidic aqueous medium. Other potentially useful matrix materials comprise collagen, homopolymers and copolymers of glycolic acid and lactic acid,

hydroxyapatite, tricalcium phosphate and other calcium phosphates.

The osteogenic proteins and implantable osteogenic devices enabled and disclosed herein will permit the physician to obtain optimal predictable bone formation to correct, for example, acquired and congenital craniofacial and other skeletal or dental anomalies (Glowacki et al. (1981) Lancet 1:959-963). The devices may be used to induce local endochondral bone formation in non-union fractures as demonstrated in animal tests, and in other clinical applications including dental and periodontal applications where bone formation is required. Another potential clinical application is in cartilage repair, for example, in the treatment of osteoarthritis.

Brief Description of the Drawing

The foregoing and other objects of this invention, the various features thereof, as well as the invention itself, may be more fully understood from the following description, when read together with the accompanying drawings, in which:

FIGURE 1 compares the amino acid sequences of the mature mOP-2 and hOP-2 polypeptide chains: hOP2-A and mOP2-A; and

25 FIGURE 2 compares the amino acid sequences of the mature OP1 and OP2 polypeptide chains: OP1-18, mOP1-S, hOP2-A and mOP2-A.

Description

Purification protocols first were developed which enabled isolation of the osteogenic protein present in crude protein extracts from mammalian bone. (See PCT WO 89/09787, published 19-OCT-89, and U.S. Serial No. 05 179,406 filed April 8, 1988, now U.S. Patent No. 4,968,950). The development of the procedure, coupled with the availability of fresh calf bone, enabled isolation of substantially pure bovine osteogenic protein (bOP). bOP was characterized significantly; 10 its ability to induce cartilage and ultimately endochondral bone growth in cat, rabbit, and rat were demonstrated and studied; it was shown to be able to induce the full developmental cascade of bone formation previously ascribed to unknown protein or proteins in 15 heterogeneous bone extracts. This dose dependent and highly specific activity was present whether or not the protein was glycosylated (see Sampath et al., (1990) J. Biol. Chem. 265: 13198-13205). Sequence data obtained from the bovine materials suggested probe designs which 20 were used to isolate human genes. The OP human counterpart proteins have now been expressed and extensively characterized.

These discoveries enabled preparation of DNAs encoding totally novel, non-native protein constructs

25 which individually as homodimers and combined with other species as heterodimers are capable of producing true endochondral bone (see PCT WO 09788, published 19-OCT-89, and US Serial No. 315,342, filed 23-FEB-89, now U.S. Patent No. 5,011,691). They also permitted expression of the natural material, truncated forms, muteins, analogs, fusion proteins, and various other variants and constructs, from cDNAs and genomic DNAs retrieved from natural sources or from synthetic DNA produced using the techniques disclosed herein and

- 13 -

using automated, commercially available equipment. The DNAs may be expressed using well established molecular biology and recombinant DNA techniques in procaryotic or eucaryotic host cells, and may be oxidized and refolded in vitro if necessary, to produce biologically active protein.

One of the DNA sequences isolated from human genomic and cDNA libraries encoded a previously unidentified gene, referred to herein as OP1. The protein encoded by the isolated DNA was identified originally by amino acid homology with proteins in the TGF-β family. Consensus splice signals were found where amino acid homologies ended, designating exonintron boundaries. Three exons were combined to obtain a functional TGF-β-like domain containing seven cysteines. (See, for example, U.S. Patent No. 5,011,691, or Ozkaynak, E. et al., (1990) EMBO. 9: 2085-2093).

The full-length cDNA sequence for hOP1, and its encoded "prepro" form "hOP1-PP," which includes an N-20 terminal signal peptide sequence, are disclosed in Seq. ID No. 1 (residues 1-431). The mature form of the hOP1 protein expressed in mammalian cells, "OP1-18", is described by amino acid residues 293-431 of Seq. ID No. 1. The full length form of hOP1, as well as 25 various truncated forms of the gene, and fused genes, have been expressed in \underline{E} . \underline{coli} and numerous mammalian cells (see, for example, published PCT application WO 91/05802, published 2-MAY-91) and all have been shown to have osteogenic activity when implanted in a mammal 30 in association with a suitable matrix.

Given the foregoing amino acid and DNA sequence information, various nucleic acids (RNAs and DNAs) can be constructed which encode at least the active region of the hOPl protein (e.g., OPS or OP7) and various of analogs thereof (including allelic and species variants and those containing genetically engineered mutations), as well as fusion proteins, truncated forms of the mature proteins, and similar constructs. Moreover, DNA hybridization probes can be constructed from fragments of the hOPl DNA or designed de novo based on the hOPl DNA or amino acid sequence. These probes then can be used to screen different genomic and cDNA libraries to identify additional osteogenic proteins.

The DNAs can be produced by those skilled in the

15 art using well known DNA manipulation techniques
involving genomic and cDNA isolation, construction of
synthetic DNA from synthesized oligonucleotides, and
cassette mutagenesis techniques. 15-100mer
oligonucleotides may be synthesized on a Biosearch DNA
20 Model 8600 Synthesizer, and purified by polyacrylamide
gel electrophoresis (PAGE) in Tris-Borate-EDTA buffer.
The DNA then may be electroeluted from the gel.
Overlapping oligomers may be phosphorylated by T4
polynucleotide kinase and ligated into larger blocks
which may also be purified by PAGE.

DNAs used as hybridization probes may be labelled (e.g., as with a radioisotope, by nick-translation) and used to identify clones in a given library containing DNA to which the probe hybridizes, following techniques well known in the art. The libraries may be obtained commercially or they may constructed de novo using conventional molecular biology techniques. Further information on DNA library construction and

hybridization techniques can be found in numerous texts known to those skilled in the art. See, for example, F.M. Ausubel., ed., <u>Current Protocols in Molecular Biology-Vol. 1</u>, (1989). In particular, see unit 5, "Construction of Recombinant DNA Libraries" and Unit 6, "Screening of Recombinant Libraries."

The DNA from appropriately identified clones then can be isolated, subcloned (preferably into an expression vector), and sequenced. Plasmids containing 10 sequences of interest then can be transfected into an appropriate host cell for protein expression and further characterization. The host may be a procaryotic or eucaryotic cell since the former's inability to glycosylate protein will not destroy the 15 protein's osteogenic activity. Useful host cells include E. coli, Saccharomyces, the insect/baculovirus cell system, myeloma cells, and various mammalian cells. The vector additionally may encode various sequences to promote correct expression of the 20 recombinant protein, including transcription promoter and termination sequences, enhancer sequences, preferred ribosome binding site sequences, preferred mRNA leader sequences, preferred signal sequences for protein secretion, and the like. The DNA sequence 25 encoding the gene of interest also may be manipulated to remove potentially inhibiting sequences or to minimize unwanted secondary structure formation. recombinant osteogenic protein also may be expressed as a fusion protein. After being translated, the protein 30 may be purified from the cells themselves or recovered from the culture medium. All biologically active protein forms comprise dimeric species joined by disulfide bonds or otherwise associated, produced by oxidizing and refolding one or more of the various

recombinant polypeptide chains within an appropriate eucaryotic cell or in vitro after expression of individual subunits. A detailed description of osteogenic protein expressed from recombinant DNA in E. coli is disclosed in U.S. Serial No. 660,162, filed 27-FEB-91, the disclosure of which incorporated by reference herein. A detailed description of osteogenic protein expressed from recombinant DNA in numerous different mammalian cells is disclosed in PCT W091/05802, also incorporated herein by reference.

Finally, in view of the disclosure made herein, and using standard methodologies known in the art, persosn skilled inthe art can raise polyclonal and monoclonal antibodies against all or part of a polypeptide chain disclosed herein, such that the antibodies are capable of binding specifically to an epitope on the polypeptide chain. Useful protocols can be found in, for example, Molecular Cloing-A Laboratory Manual (Sambrook et al. eds., Cold Spring Harbor Press 2nd ed. 1989). See Book 3, Section 18.

Exemplification

In an effort to identify additional DNA sequences encoding osteogenic proteins, a hybridization probe specific to the C-terminus of the DNA of mature OP-1
25 was prepared using a StuI-EcoRl digest fragment of OP-1 (base pairs 1034-1354 in Sequence ID No. 1), and labelled with ³²P by nick translation, as described in the art. As disclosed supra, the OP1 C-terminus encodes a key functional domain, e.g., the "active region" for osteogenic activity. The C-terminus also is the region of the protein whose amino acid sequence shares specific amino acid sequence homology with

particular proteins in the TGF- β super-family of regulatory proteins, and which includes the conserved cysteine skeleton.

Approximately 7 x 10⁵ phages of an oligo(dT) primed 17.5 days p.c. mouse embryo 5' stretch cDNA (gt10) library (Clonetech, Inc., Palo Alto, CA) was screened with the labelled probe. The screen was performed using the following stringent hybridization conditions: 40% formamide, 5 x SSPE, 5 x Denhart's solution, 0.1% SDS, at 37°C overnight, and washing in 0.1 x SSPE, 0.1% SDS at 50°C.

Five recombinant phages were purified over three rounds of screening. Phage DNA was prepared from all five phages, subjected to an EcoRl digest, subcloned into the EcoRl site of a common puC-type plasmid modified to allow single strand sequencing, and sequenced using means well known in the art.

Two different DNAs were identified by this procedure. One DNA, referred to herein as mOP1, has substantial homology to the mature form of OP1 (about 98%), and is described in detail in copending USSN 600,024, filed 18-Oct-90. A second DNA, encoding the C-terminus of a related gene and referred to herein as mOP2, also was identified by this procedure. The N-terminus of the gene encoding mOP2 was identified subsequently by screening a second mouse cDNA library (Mouse PCC4 cDNA (ZAP) library, Stratagene, Inc., La Jolla, CA).

Mouse OP2 (mOP2) protein shares significant amino acid sequence homology with the amino acid sequence of the hOP1 active region, e.g., OPS or OP7, about 74%

homology, and less homology with the intact mature form, e.g., OP1-18, about 58% homology. The cDNA sequence, and the encoded amino acid sequence, for the full length mOP-2 protein is depicted in Sequence ID 05 No. 3. The full-length form of the protein is referred to as the prepro form of mOP-2 ("mOP2-PP"), and includes a signal peptide sequence at its N-terminus. The amino acid sequence Leu-Ala-Leu-Cys-Ala-Leu (amino acid residues 13-18 of Sequence ID No. 3) is believed 10 to constitute the cleavage site for the removal of the signal peptide sequence, leaving an intermediate form of the protein, the "pro" form, to be secreted from the expressing cell. The amino acid sequence Arg-Ala-Pro-Arg-Ala (amino acid residues 255-259 of Sequence ID 15 No. 3) is believed to constitute the cleavage site that produces the mature form of the protein, herein referred to as "mOP2-A", and described by residues 259-397 of Seq. ID No. 3. Residues 301-397 of Seq. ID No. 3 correspond to the region defining the conserved $_{20}$ six cysteine skeleton. Residues 296-397 of Seq. ID No. 3 correspond to the region defining the conserved seven cysteine skeleton.

Using a probe prepared from the pro region of mOP2 (an EcoRl-BamH1 digest fragment, bp 467-771 of Sequence ID No. 3), a human hippocampus library was screened (human hippocampus cDNA lambda (ZAP II library Stratagene, Inc., La Jolla, CA) following essentially the same procedure as for the mouse library screens. The procedure identified the N-terminus of a novel DNA encoding an amino acid sequence having substantial homology with mOP2. The C-terminus of the gene subsequently was identified by probing a human genomic library (in lambda phage EMBL-3, Clonetech, Inc., Palo Alto, CA) with a labelled fragment from the novel human

DNA in hand. The novel polypeptide chain encoded by this DNA is referred to herein as hOP2 protein, and shares almost complete amino acid identity (about 92% amino acid sequence homology) with mOP2-A (see Fig. 1 and infra).

05

25

30

The cDNA sequence, and the encoded amino acid sequence, for the prepro form of hOP2, "hOP2-PP", is described in Sequence ID No. 5. This full-length form of the protein also includes a signal peptide sequence at its N-terminus. The amino acid sequence Leu-Ala-10 Leu-Cys-Ala-Leu (amino acid residues 13-18 of Sequence ID No. 5) is believed to constitute the cleavage site for the removal of the signal peptide sequence, leaving an intermediate form of the protein, the "pro" form, to be secreted from the expressing cell. The amino acid 15 sequence Arg-Thr-Pro-Arg-Ala (amino acid residues 257-261 of Sequence ID No. 5) is believed to constitute the cleavage site that produces what is believed to be the mature form of the protein, herein referred to as hOP2-A" and described by residues 261-399 of Seq. ID 20 No. 5.

Additional mature species of hOP2 thought to be active include truncated sequences, "hOP2-P" (described by residues 264-399 of Seq. ID No. 5) and "hOP2-R" (described by residues 267-399 of Seq. ID No. 5), and a slightly longer sequence ("hOP2-S", described by residues 240-399 of Seq. ID No. 5). Residues 303-399 of Seq. ID No. 5 correspond to the region defining the conserved six cysteine skeleton. Residues 297-399 of Seq. ID No. 5 correspond to the region defining the conserved seven cystein skeleton.

It should be noted that the nucleic acid sequence encoding the N-terminus of the prepro form of both mOP2 and hOP2 is rich in guanidine and cytosine base pairs. As will be appreciated by those skilled in the art, 05 sequencing such a "G-C rich" region can be problematic, due to stutter and/or band compression. Accordingly, the possibility of sequencing errors in this region can not be ruled out. However, the definitive amino acid sequence for these and other, similarly identified proteins can be determined readily by expressing the protein from recombinant DNA using, for example, any of the means disclosed herein, and sequencing the polypeptide chain by conventional peptide sequencing methods well known in the art.

15 Figure 1 compares the amino acid sequences of mature mOP2 and hOP2. Identity is indicated by three dots (...) in the mOP2 sequence. As is evident from the figure, the amino acid sequence homology between the mature forms of these two proteins is substantial 20 (92% homology between the mature sequences, about 95% homology within the C-terminal active region (e.g., residues 38-139 or 42-139 of Fig. 1.)

Fig. 2 compares the amino acid sequences for the mature forms of all four species of OP1 and OP2 proteins. Here again, identity is indicated by three dots (...). Like the mOP2 protein, the hOP2 protein shares significant homology (about 74%) with the amino acid sequence defining the OP1 active region (OPS or OP7, residues 43-139 and 38-139, respectively, in Fig. 2), and less homology with OP1-18 (about 58% homology). Both OP2 proteins share the conserved seven cysteine skeleton seen in the OP1 proteins. In

addition, the OP2 proteins comprise an eighth cysteine residue within this region (see position 78 in FIG. 2).

A preferred generic amino acid sequence useful as a subunit of a dimeric osteogenic protein capable of inducing endochondral bone or cartilage formation when implanted in a mammal in association with a matrix, and which incorporates the maximum homology between the identified OP1 and OP2 proteins, can be described by the sequence referred to herein as "OPX", described below and in Seq. No.7.

	Cys	Xaa	Xaa	His	Glu	Leu	Tyr	Val	Xaa	Phe
	1				5					10
15	Xaa	Asp	Leu	Gly	Trp	Xaa	Asp	Trp	Xaa	Ile
					15					20
	Ala	Pro	Xaa	Gly	Tyr	Xaa	Ala	Tyr	Tyr	Cys
					25					30
	Glu	Gly	Glu	Cys	Xaa	Phe	Pro	Leu	Xaa	
20					35					40
	Xaa	Met	Asn	Ala	Thr	Asn	His	Ala	Ile	
					45					50
	Gln	Xaa	Leu	Val	His	Xaa	Xaa	Xaa	Pro	
					55					60
25	Xaa	Val	Pro	Lys	Xaa	Cys	Cys	Ala	Pro	
					65					70
	Xaa	Leu	Xaa	Ala	Xaa	Ser	Val	Leu	Tyr	
					75					80
	Asp	Xaa	Ser	Xaa	Asn	Val	Xaa	Leu	Xaa	
30					85					90
	Xaa	Arg	Asn	Met	Val	Val	Xaa	Ala	Cys	
					95					100
	Cys	His	,							

and wherein Xaa at res. 2 = (Lys or Arg); Xaa at res. 3 = (Lys or Arg); Xaa at res. 9 = (Ser or Arg); Xaa at res. 11 = (Arg or Gln); Xaa at res. 16 = (Gln or Leu); Xaa at res. 19 = (Ile or Val); Xaa at res. 23 = 5 (Glu or Gln); Xaa at res. 26 = (Ala or Ser); Xaa at res. 35 = (Ala or Ser); Xaa at res. 39 = (Asn or Asp); Xaa at res. 41 = (Tyr or Cys); Xaa at res. 50 = (Val or Leu); Xaa at res. 52 = (Ser or Thr); Xaa at res. 56 = (Phe or Leu); Xaa at res. 57 = (Ile or Met); Xaa at 10 res. 58 = (Asn or Lys); Xaa at res. 60 = (Glu, Asp or Asn); Xaa at res. 61 = (Thr, Ala or Val); Xaa at res. 65 = (Pro or Ala); Xaa at res. 71 = (Gln or Lys); Xaa at res. 73 = (Asn or Ser); Xaa at res. 75 = (Ile orThr); Xaa at res. 80 = (Phe or Tyr); Xaa at res. 82 = 15 (Asp or Ser); Xaa at res. 84 = (Ser or Asn); Xaa at res. 87 = (Ile or Asp); Xaa at res. 89 = (Lys or Arg); Xaa at res. 91 = (Tyr, Ala or His); and Xaa at res. 97 = (Arg or Lys).

The high degree of homology exhibited between the 20 various OP1 and OP2 proteins suggests that the novel osteogenic proteins identified herein will purify essentially as OP1 does, or with only minor modifications of the protocols disclosed for OP1. 25 Similarly, the purified mOP1, mOP2, and hOP2 proteins are predicted to have an apparent molecular weight of about 18 kDa as reduced single subunits, and an apparent molecular weight of about 36 kDa as oxidized dimers, as determined by comparison with molecular 30 weight standards on an SDS-polyacrylamide electrophoresis gel. Unglycosylated dimers (e.g., proteins produced by recombinant expression in E. coli) are predicted to have an apparent molecular weight of about 27 kDa. There appears to be one potential N

glycosylation site in the mature forms of the mOP2 and hOP2 proteins.

The identification of osteogenic proteins having an active region comprising eight cysteine residues also allows one to construct osteogenic polypeptide chains patterned after either of the following template amino 05 acid sequences, or to identify additional osteogenic proteins having this sequence. The template sequences contemplated are "OPX-7C", comprising the conserved six 10 cysteine skeleton plus the additional cysteine residue identified in the OP2 proteins, and "OPX-8C", comprising the conserved seven cysteine skeleton plus the additional cysteine residue identified in the OP2 proteins. The OPX-7C and OPX-8C sequences are described below and in Seq. ID Nos. 8 and 9, respectively. Each Xaa in these template sequences independently represents one of the 20 naturallyoccurring L-isomer, α -amino acids, or a derivative thereof. Biosynthetic constructs patterned after this template readily are constructed using conventional DNA 20 synthesis or peptide synthesis techniques well known in the art. Once constructed, osteogenic proteins comprising these polypeptide chains can be tested as disclosed herein.

25 "OPX-7C" (Sequence ID No. 8):

 Xaa
 X

		2.5					40				
	Vaa	35 Xaa X	(aa)	Kaa	Xaa	Xaa	Xaa	Xaa	Xaa	Xaa	Xaa
	4.5					50					
	Xaa	Xaa >	(aa)	Xaa	Xaa	Cys	Cys	Xaa	Xaa	Xaa 65	Хаа
05					60					00	
	Xaa	Xaa 2	Xaa :	Хаа 70	хаа	Xaa	You	Auu	75		
	Vaa	Xaa 2	Хаа	Xaa	Xaa	Xaa	Xaa	Xaa	Xaa	Xaa	Xaa
			80					85			
10	Xaa	Xaa	Xaa	Xaa	Xaa	Cys	Xaa	Cys	Xaa		
		90					95				
"OPX-8C" (Sequence ID No. 9 comprising additional five									l five		
"0	PX-8C" (Se esidues at	quenc the N	e 11 I-te:	cmin	us,	incl	udir	ıg a	cons	erve	d.
re	steine res	idue)) :								
C,							W- -	Vas	Yaa	Xaa	
15	Cys Xaa	Xaa >	Kaa :	Xaa	Xaa	хаа	хаа	Add	10		
	1 Xaa Xaa	Vaa '	Vaa	5 Xaa	Xaa	Xaa	Xaa	Xaa	Xaa	Xaa	
			1.5					20			
	Xaa Xaa	Xaa	Xaa	Xaa	Xaa	Xaa	Cys	Xaa	Xaa	Xaa	
20		2 5					30				Xaa
	Cys Xaa		Xaa	Xaa	хаа	xaa 40	Cys)	nuu	•••		45
	35 Xaa Xaa	Yaa	Xaa	Xaa	Xaa			xaa	Xaa	Хаа	L
				50						•	
25	Xaa Xaa	Xaa	Xaa	Xaa	Xaa	xaa	a Xaa	a Xaa 6!	a Cys	5	
			60	••	Va	. Va	a Xa			a Xaa	a
	Cys Xaa	a Xaa	Xaa 70		, Aac	ı Au	4.00	7	5		
	Xaa Xa	a Xaa	Xaa	Xaa	a Xa	a Xa	a Xa	a Xa	a Xa	a Xa	a
00		80					0	5			
30	Xaa Xa	a Xaa	Xaa	a Xa	a Xa	a Xa	a Xa	a Xa	a xa	a Cy	5
		0				g	5				
	хаа Су	s Xaa	3								

- 25 -

100

MATRIX PREPARATION

A. General Consideration of Matrix Properties

The currently preferred carrier material is a xenogenic bone-derived particulate matrix treated as disclosed herein. This carrier may be replaced by either a biodegradable-synthetic or synthetic-inorganic matrix (e.g., hydroxylapatite (HAP), collagen, tricalcium phosphate or polylactic acid, polyglycolic acid and various copolymers thereof.)

Studies have shown that surface charge, particle size, the presence of mineral, and the methodology for combining matrix and osteogenic protein all play a role in achieving successful bone induction. Perturbation of the charge by chemical modification abolishes the inductive response. Particle size influences the quantitative response of new bone; particles between 75 µm and 420 µm elicit the maximum response. Contamination of the matrix with bone mineral will inhibit bone formation. Most importantly, the procedures used to formulate OP onto the matrix are extremely sensitive to the physical and chemical state of both the osteogenic protein and the matrix.

The sequential cellular reactions in the
interface of the bone matrix/osteogenic protein
implants are complex. The multistep cascade includes:
binding of fibrin and fibronectin to implated matrix,
chemotaxis of cells, proliferation of fibroblasts,
differentiation into chondroblasts, cartilage

PCT/US91/07635 WO 92/07073

formation, vascular invasion, bone formation, remodeling, and bone marrow differentiation.

A successful carrier for osteogenic protein must perform several important functions. It must bind osteogenic protein and act as a slow release delivery 05 system, accommodate each step of the cellular response during bone development, and protect the osteogenic protein from nonspecific proteolysis. In addition, selected materials must be biocompatible in vivo and preferably biodegradable; the carrier must act as a temporary scaffold until replaced completely by new 10 bone. Polylactic acid (PLA), polyglycolic acid (PGA), and various combinations have different dissolution rates in vivo. In bones, the dissolution rates can vary according to whether the implant is placed in 15 cortical or trabecular bone.

Matrix geometry, particle size, the presence of surface charge, and the degree of both intra-andinter-particle porosity are all important to successful matrix performance. It is preferred to shape the matrix to the desired form of the new bone and to have dimensions which span non-union defects. Rat studies show that the new bone is formed essentially having the dimensions of the device implanted.

20

25

The matrix may comprise a shape-retaining solid made of loosely adhered particulate material, e.g., with collagen. It may also comprise a molded, porous solid, or simply an aggregation of close-packed particles held in place by surrounding tissue. Masticated muscle or other tissue may also be used. 30 Large allogenic bone implants can act as a carrier for the matrix if their marrow cavities are cleaned and

packed with particle and the dispersed osteogenic protein.

The preferred matrix material, prepared from xenogenic bone and treated as disclosed herein, 05 produces an implantable material useful in a variety of clinical settings. In addition to its use as a matrix for bone formation in various orthopedic, periodontal, and reconstructive procedures, the matrix also may be used as a sustained release carrier, or as a 10 collagenous coating for implants. The matrix may be shaped as desired in anticipation of surgery or shaped by the physician or technician during surgery. the material may be used for topical, subcutaneous, intraperitoneal, or intramuscular implants; it may be 15 shaped to span a nonunion fracture or to fill a bone defect. In bone formation or conduction procedures, the material is slowly absorbed by the body and is replaced by bone in the shape of or very nearly the shape of the implant.

Various growth factors, hormones, enzymes, therapeutic compositions, antibiotics, and other body treating agents also may be absorbed onto the carrier material and will be released over time when implanted as the matrix material is slowly absorbed. Thus, various known growth factors such as EGF, PDGF, IGF, FGF, TGF-α, and TGF-β may be released in vivo. The material can be used to release chemotherapeutic agents, insulin, enzymes, or enzyme inhibitors.

B. Bone-Derived Matrices

Preparation of Demineralized Bone

Demineralized bone matrix, preferably bovine bone matrix, is prepared by previously published procedures (Sampath and Reddi (1983) Proc. Natl. Acad. Sci. USA 80:6591-6595). Bovine diaphyseal bones (age 05 1-10 days) are obtained from a local slaughterhouse and used fresh. The bones are stripped of muscle and fat, cleaned of periosteum, demarrowed by pressure with cold water, dipped in cold absolute ethanol, and stored at -20°C. They are then dried and fragmented by crushing 10 and pulverized in a large mill. Care is taken to prevent heating by using liquid nitrogen. pulverized bone is milled to a particle size in the range of 70-850 μ m, preferably 150-420 μ m, and is defatted by two washes of approximately two hours 15 duration with three volumes of chloroform and methanol (3:1). The particulate bone is then washed with one volume of absolute ethanol and dried over one volume of anhydrous ether yielding defatted bone powder. defatted bone powder is then demineralized by four 20 successive treatments with 10 volumes of 0.5 N HCl at 4°C for 40 min. Finally, neutralizing washes are done on the demineralized bone powder with a large volume of water.

2. Guanidine Extraction

Demineralized bone matrix thus prepared is extracted with 5 volumes of 4 M guanidine-HCl, 50mM Tris-HCl, pH 7.0 for 16 hr. at 4°C. The suspension is filtered. The insoluble material is collected and used to fabricate the matrix. The material is mostly collagenous in nature. It is devoid of osteogenic or chondrogenic activity.

3. Matrix Treatments

The major component of all bone matrices is Type-I collagen. In addition to collagen, demineralized bone extracted as disclosed above 05 includes non-collagenous proteins which may account for 5% of its mass. In a xenogenic matrix, these noncollagenous components may present themselves as potent antigens, and may constitute immunogenic and/or inhibitory components. These components also may 10 inhibit osteogenesis in allogenic implants by interfering with the developmental cascade of bone differentiation. It has been discovered that treatment of the matrix particles with a collagen fibril-modifying agent extracts potentially unwanted 15 components from the matrix, and alters the surface structure of the matrix material. Useful agents include acids, organic solvents or heated aqueous media. Various treatments are described below. A detailed physical analysis of the effect these fibril-20 modifying agents have on demineralized, quanidineextracted bone collagen particles is disclosed in PCT WO 90/10018, published 7-SEP-90.

After contact with the fibril-modifying agent, the treated matrix is washed to remove any extracted components, following a form of the procedure set forth below:

Suspend in TBS (Tris-buffered saline)
 1g/200 ml and stir at 4°C for 2 hrs; or in 6 M urea, 50 mM Tris-HCl, 500 mM NaCl, pH 7.0 (UTBS) or water and
 stir at room temperature (RT) for 30 minutes (sufficient time to neutralize the pH);

- 30 -

- 2. Centrifuge and repeat wash step; and
- 3. Centrifuge; discard supernatant; water wash residue; and then lyophilize.

3.1 Acid Treatments

05 1. Trifluoroacetic acid.

Trifluoroacetic acid is a strong non-oxidizing acid that is a known swelling agent for proteins, and which modifies collagen fibrils.

Bovine bone residue prepared as described

10 above is sieved, and particles of the appropriate size are collected. These particles are extracted with various percentages (1.0% to 100%) of trifluoroacetic acid and water (v/v) at 0°C or room temperature for 1-2 hours with constant stirring. The treated matrix is

15 filtered, lyophilized, or washed with water/salt and then lyophilized.

2. Hydrogen Fluoride.

Like trifluoroacetic acid, hydrogen fluoride is a strong acid and swelling agent, and also is 20 capable of altering intraparticle surface structure. Hydrogen fluoride is also a known deglycosylating agent. As such, HF may function to increase the osteogenic activity of these matrices by removing the antigenic carbohydrate content of any glycoproteins 25 still associated with the matrix after guanidine extraction.

Bovine bone residue prepared as described above is sieved, and particles of the appropriate size are collected. The sample is dried in vacuo over P205, transferred to the reaction vessel and exposed to 05 anhydrous hydrogen fluoride (10-20 ml/g of matrix) by distillation onto the sample at -70°C. The vessel is allowed to warm to 0°C and the reaction mixture is stirred at this temperature for 120 minutes. After evaporation of the hydrogen fluoride in vacuo, the 10 residue is dried thoroughly in vacuo over KOH pellets to remove any remaining traces of acid. Extent of deglycosylation can be determined from carbohydrate analysis of matrix samples taken before and after treatment with hydrogen fluoride, after washing the 15 samples appropriately to remove non-covalently bound carbohydrates. SDS-extracted protein from HF-treated material is negative for carbohydrate as determined by Con A blotting.

The deglycosylated bone matrix is next washed 20 twice in TBS (Tris-buffered saline) or UTBS, water-washed, and then lyophilized.

Other acid treatments are envisioned in addition to HF and TFA. TFA is a currently preferred acidifying reagent in these treatments because of its 25 volatility. However, it is understood that other, potentially less caustic acids may be used, such as acetic or formic acid.

3.2 Solvent Treatment

1. Dichloromethane.

Dichloromethane (DCM) is an organic solvent capable of denaturing proteins without affecting their primary structure. This swelling agent is a common reagent in automated peptide synthesis, and is used in 05 washing steps to remove components.

Bovine bone residue, prepared as described above, is sieved, and particles of the appropriate size are incubated in 100% DCM or, preferably, 99.9% DCM/0.1% TFA. The matrix is incubated with the 10 swelling agent for one or two hours at 0°C or at room temperature. Alternatively, the matrix is treated with the agent at least three times with short washes (20 minutes each) with no incubation.

Acetonitrile. 2.

Acetonitrile (ACN) is an organic solvent, 15 capable of denaturing proteins without affecting their primary structure. It is a common reagent used in high-performance liquid chromatography, and is used to elute proteins from silica-based columns by perturbing 20 hydrophobic interactions.

Bovine bone residue particles of the appropriate size, prepared as described above, are treated with 100% ACN (1.0 g/30 ml) or, preferably, 99.9% ACN/0.1% TFA at room temperature for 1-2 hours 25 with constant stirring. The treated matrix is then water-washed, or washed with urea buffer, or 4 M NaCl and lyophilized. Alternatively, the ACN or ACN/TFA treated matrix may be lyophilized without wash.

3. Isopropanol.

Isopropanol is also an organic solvent capable of denaturing proteins without affecting their primary structure. It is a common reagent used to elute proteins from silica HPLC columns.

Bovine bone residue particles of the appropriate size prepared as described above are treated with 100% isopropanol (1.0 g/30 ml) or, preferably, in the presence of 0.1% TFA, at room temperature for 1-2 hours with constant stirring. The matrix is then water-washed or washed with urea buffer or 4 M NaCl before being lyophilized.

4. Chloroform

Chloroform also may be used to increase surface area of bone matrix like the reagents set forth above, either alone or acidified.

Treatment as set forth above is effective to assure that the material is free of pathogens prior to implantation.

3.3 Heat Treatment

The currently most preferred agent is a heated aqueous fibril-modifying medium such as water, to increase the matrix particle surface area and porosity. The currently most preferred aqueous medium is an acidic aqueous medium having a pH of less than about 4.5, e.g., within the range of pH 2 - pH 4. which may help to "swell" the collagen before heating. 0.1% acetic acid, which has a pH of about 3, currently is preferred. O.1 M acetic acid also may be used.

Various amounts of delipidated, demineralized guanidine-extracted bone collagen are heated in the aqueous medium (1g matrix/30ml aqueous medium) under constant stirring in a water jacketed glass flask, and 05 maintained at a given temperature for a predetermined period of time. Preferred treatment times are about one hour, although exposure times of between about 0.5 to two hours appear acceptable. The temperature employed is held constant at a temperature generally within the range of about 37°C to 75°C. The currently preferred heat treatment temperature is within the range of 45°C to 60°C.

After the heat treatment, the matrix is filtered, washed, lyophilized and used for implant. Where an acidic aqueous medium is used, the matrix also is preferably neutralized prior to washing and lyophilization. A currently preferred neutralization buffer is a 200mM sodium phosphate buffer, pH 7.0. To neutralize the matrix, the matrix preferably first is allowed to cool following thermal treatment, the acidic aqueous medium (e.g., 0.1% acetic acid) then is removed and replaced with the neutralization buffer and the matrix agitated for about 30 minutes. The neutralization buffer then may be removed and the matrix washed and lyophilized (see infra).

The matrix also may be treated to remove contaminating heavy metals, such as by exposing the matrix to a metal ion chelator. For example, following thermal treatment with 0.1% acetic acid, the matrix may be neutralized in a neutralization buffer containing EDTA (sodium ethylenediaminetetraacetic acid), e.g., 200 mM sodium phosphate, 5mM EDTA, pH 7.0. 5 mM EDTA provides about a 100-fold molar excess of chelator to

residual heavy metals present in the most contaminated matrix tested to date. Subsequent washing of the matrix following neutralization appears to remove the bulk of the EDTA. EDTA treatment of matrix particles reduces the residual heavy metal content of all metals tested (Sb, As, Be, Cd, Cr, Cu, Co, Pb, Hg, Ni, Se, Ag, Zn, Tl) to less than about 1 ppm. Bioassays with EDTA-treated matrices indicate that treatment with the metal ion chelator does not inhibit bone inducing activity.

The collagen matrix materials preferably take the form of a fine powder, insoluble in water, comprising nonadherent particles. It may be used simply by packing into the volume where new bone growth or sustained release is desired, held in place by surrounding tissue. Alternatively, the powder may be encapsulated in, e.g., a gelatin or polylactic acid coating, which is adsorbed readily by the body. The powder may be shaped to a volume of given dimensions and held in that shape by interadhering the particles using, for example, soluble, species-biocompatible collagen. The material may also be produced in sheet, rod, bead, or other macroscopic shapes.

Demineralized rat bone matrix used as an allogenic matrix in certain of the experiments

25 disclosed herein, is prepared from several of the dehydrated diaphyseal shafts of rat femur and tibia as described herein to produce a bone particle size which passes through a 420 μ m sieve. The bone particles are subjected to dissociative extraction with 4 M guanidine-HCl. Such treatment results in a complete loss of the inherent ability of the bone matrix to induce endochondral bone differentiation. The remaining insoluble material is used to fabricate the

WO 92/07073 PCT/US91/07635

- 36 -

matrix. The material is mostly collagenous in nature, and upon implantation, does not induce cartilage and bone. All new preparations are tested for mineral content and osteogenic activity before use. The total loss of biological activity of bone matrix is restored when an active osteoinductive protein fraction or a pure osteoinductive protein preparation is reconstituted with the biologically inactive insoluble collagenous matrix.

FABRICATION OF OSTEOGENIC DEVICE

The naturally sourced and recombinant protein as set forth above, and other constructs, can be combined and dispersed in a suitable matrix preparation using any of the methods described below. In general, 50-100 ng of active protein is combined with the inactive carrier matrix (e.g., 25 mg for rat bioassays). Greater amounts may be used for large implants.

Ethanol Precipitation

10

Matrix is added to osteogenic protein

20 dissolved in guanidine-HCl. Samples are vortexed and incubated at a low temperature (e.g., 4°C). Samples are then further vortexed. Cold absolute ethanol (5 volumes) is added to the mixture which is then stirred and incubated, preferably for 30 minutes at -20°C.

25 After centrifugation (microfuge, high speed) the supernatant is discarded. The reconstituted matrix is washed twice with cold concentrated ethanol in water (85% EtOH) and then lyophilized.

Acetonitrile Trifluoroacetic

Acid Lyophilization

In this procedure, osteogenic protein in an acetonitrile trifluroacetic acid (ACN/TFA) solution is added to the carrier material. Samples are vigorously vortexed many times and then lyophilized. This method is currently preferred, and has been tested with osteogenic protein at varying concentrations and different levels of purity.

Urea Lyophilization

For those osteogenic proteins that are prepared in urea buffer, the protein is mixed with the matrix material, vortexed many times, and then lyophilized. The lyophilized material may be used "as is" for implants.

15 4. Buffered Saline Lyophilization

OP1 and OP2 preparations in physiological saline may also be vortexed with the matrix and lyophilized to produce osteogenically active material.

These procedures also can be used to adsorb other active therapeutic drugs, hormones, and various bioactive species to the matrix for sustained release purposes.

BIOASSAY

The functioning of the various proteins and

25 devices of this invention can be evaluated with an <u>in</u>

<u>vivo</u> bioassay. Studies in rats show the osteogenic

effect in an appropriate matrix to be dependent on the

dose of osteogenic protein dispersed in the matrix. No activity is observed if the matrix is implanted alone.

In vivo bioassays performed in the rat model also have shown that demineralized, guanidine-extracted xenogenic bone matrix materials of the type described in the literature are ineffective as a carrier, fail to induce bone, and produce an inflammatory and immunological response when implanted unless treated as disclosed above. In certain species (e.g., monkey) allogenic matrix materials also apparently are ineffective as carriers. The following sets forth various procedures for preparing osteogenic devices from the proteins and matrix materials prepared as set forth above, and for evaluating their osteogenic utility.

15 A. Rat Model

1. Implantation

The bioassay for bone induction as described by Sampath and Reddi ((1983) Proc. Natl. Acad. Sci. USA 80 6591-6595), herein incorporated by reference, may be 20 used to monitor endochondral bone differentiation activity. This assay consists of implanting test samples in subcutaneous sites in recipient rats under ether anesthesia. Male Long-Evans rats, aged 28-32 days, were used. A vertical incision (1 cm) is made 25 under sterile conditions in the skin over the thoracic region, and a pocket is prepared by blunt dissection. Approximately 25 mg of the test sample is implanted deep into the pocket and the incision is closed with a metallic skin clip. The day of implantation is 30 designated as day one of the experiment. Implants were removed on day 12. The heterotropic site allows for the study of bone induction without the possible

ambiguities resulting from the use of orthotropic sites. As disclosed herein, both allogenic (rat bone matrix) and xenogenic (bovine bone matrix) implants were assayed.

05 2. Cellular Events

Successful implants exhibit a controlled progression through the stages of protein-induced endochondral bone development, including: (1) transient infiltration by polymorphonuclear leukocytes on day 10 one; (2) mesenchymal cell migration and proliferation on days two and three; (3) chondrocyte appearance on days five and six; (4) cartilage matrix formation on day seven; (5) cartilage calcification on day eight; (6) vascular invasion, appearance of osteoblasts, and 15 formation of new bone on days nine and ten; (7) appearance of osteoblastic and bone remodeling and dissolution of the implanted matrix on days twelve to eighteen; and (8) hematopoietic bone marrow differentiation in the ossicle on day twenty-one. 20 results show that the shape of the new bone conforms to the shape of the implanted matrix.

3. Histological Evaluation

Histological sectioning and staining is preferred to determine the extent of osteogenesis in the implants. Implants are fixed in Bouins Solution, embedded in paraffin, and cut into 6-8 μ m sections. Staining with toluidine blue or hemotoxylin/eosin demonstrates clearly the ultimate development of endochondral bone. Twelve day implants are usually sufficient to determine whether the implants contain newly induced bone.

4. Biological Markers

Alkaline phosphatase activity may be used as a marker for osteogenesis. The enzyme activity may be determined spectrophotometrically after homogenization of the implant. The activity peaks at 9-10 days in vivo and thereafter slowly declines. Implants showing no bone development by histology have little or no alkaline phosphatase activity under these assay conditions. The assay is useful for quantitation and obtaining an estimate of bone formation quickly after the implants are removed from the rat. Alternatively, the amount of bone formation can be determined by measuring the calcium content of the implant.

The invention may be embodied in other specific

15 forms without departing from the spirit or essential characteristics thereof. The present embodiments are therefore to be considered in all respects as illustrative and not restrictive, the scope of the invention being indicated by the appended claims rather than by the foregoing description, and all changes which come within the meaning and range of equivalency of the claims are therefore intended to be embraced therein.

SEQUENCE LISTING

		
(1)	GENERAL	INFORMATION:
05	(i)	APPLICANT: OPPERHANN, HERMANN OZKAYNAK, ENGIN KUBERASAMPATH, THANGAVEL RUEGER, DAVID C.
	(ii)	TITLE OF INVENTION: OSTEOGENIC DEVICES
	(iii)	NUMBER OF SEQUENCES: 9
10	(iv)	CORRESPONDENCE ADDRESS: (A) ADDRESSEE: TESTA, HURWITZ & THIBEAULT (B) STREET: 53 STATE STREET (C) CITY: BOSTON (D) STATE: MASSACHUSETTS
15		(E) COUNTRY: U.S.A. (F) ZIP: 02109
20	(▽)	COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
	(Vi)	CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: (B) FILING DATE: (C) CLASSIFICATION:
25	(viii)	ATTORNEY/AGENT INFORMATION: (A) NAME: PITCHER, EDMUND R. (B) REGISTRATION NUMBER: 27,829 (C) REFERENCE/DOCKET NUMBER: CRR056PC
30	(ix)	TELECOMMUNICATION INFORMATION: (A) TELEPHONE: 617/248-7000 (B) TELEFAX: 617/248-7100
(2)	INFORMA	TION FOR SEQ ID NO:1:
35	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 1822 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

			(4)	A NT	rı-sı	NCF.	. NO										
			(iv)					_									
			(v i)	(A	IGINA) OI) T	RGAN:	ISM:	HOH	O SA	PIEN OCAM	S PUS					٠	
	5 0		(ix)	(A (B) L) II	AME/I OCAT: DENT: THER	ION: IFIC INF /pro /evi	49. ATIO ORMA duct denc	N ME TION = "h e= E	THOD : /f OP1- XPER	unct	ion= TAL	ment "OS	al TEOG	ENIC	PROTEIN"	
			(xi)	SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:1:					
1	GGTG 5	Het His Val 1 TOA OTG CGA GCT GCG GCG CCG CAC AGC TTC GTG GCG CTC TGG GCA 105															
	CGC Arg	GTGCGGGCC CGGAGCCCGG AGCCCGGGTA GCGCGTAGAG CCGGCGCG ATG CAC GTG Het His Val 1 GC TCA CTG CGA GCT GCG GCG CCG CAC AGC TTC GTG GCG CTC TGG GCA TG Ser Leu Arg Ala Ala Ala Pro His Ser Phe Val Ala Leu Trp Ala 5 CC CTG TTC CTG CTG CGC TCC GCC CTG GCC GAC TTC AGC CTG GAC AAC TO Leu Phe Leu Leu Arg Ser Ala Leu Ala Asp Phe Ser Leu Asp Asn 25															105
20	CCC Pro 20	CTG Leu	TTC Phe	CTG Leu	CTG Leu	Arg	TCC Ser	GCC Ala	CTG Leu	GCC Ala	Asp	TTC Phe	AGC Ser	CTG Leu	GAC Asp	Asn	153
25	GAG Glu	GTG Val	CAC His	TCG Ser	AGC Ser 40	TTC Phe	ATC Ile	CAC His	CGG Arg	CGC Arg 45	CTC Leu	CGC Arg	AGC Ser	CAG Gln	GAG Glu 50	CGG Arg	201
	CGG Arg	GAG Glu	ATG Met	CAG Gln 55	CGC Arg	GAG Glu	ATC Ile	CTC Leu	TCC Ser 60	ATT Ile	TTG Leu	GGC Gly	TTG Leu	CCC Pro 65	CAC His	CGC Arg	249
30	CCG Pro	CGC Arg	CCG Pro 70	His	CTC Leu	CAG Gln	GGC Gly	AAG Lys 75	CAC	AAC Asn	TCG Ser	GCA Ala	CCC Pro 80	ATG Met	TTC Phe	ATG Met	297
	CTG Leu	GAC Asp 85	Leu	TAC	AAC Asn	GCC Ala	ATG Met 90	GCG Ala	GTG Val	GAG Glu	GAG Glu	GGC Gly 95	GGC Gly	GGG Gly	CCC Pro	GGC Gly	345
35	GGC Gly 100	Gln	GGC Gly	TTC Phe	TCC Ser	TAC Tyr 105	CCC Pro	TAC Tyr	AAG Lys	GCC	GTC Val 110	Phe	AGT Ser	ACC	CAG Gln	GGC Gly 115	3 93
	CCC Pro	CCI	CTG	GCC Ala	AGC Ser	CTG Leu	CAA Gln	GAT Asp	AGC Ser	CAT His	TTC Phe	CTC Leu	ACC	GAC Asp	GCC	GAC Asp	441

					120					125					130		
	ATG Met	GTC Val	Het	AGC Ser .35	TTC Phe	GTC Val	AAC Asn	CTC Leu	GTG Val 140	GAA Glu	CAT His	GAC Asp	AAG Lys	GAA Glu 145	TTC Phe	TTC Phe	489
05	CAC His	CCA Pro	CGC Arg 150	TAC Tyr	CAC His	CAT His	CGA Arg	GAG Glu 155	TTC Phe	CGG Arg	TTT Phe	GAT Asp	CTT Leu 160	TCC Ser	AAG Lys	ATC Ile	537
10	CCA Pro	GAA Glu 165	GGG Gly	GAA Glu	GCT Ala	GTC Val	ACG Thr 170	GCA Ala	GCC Ala	GAA Glu	TTC Phe	CGG Arg 175	ATC Ile	TAC Tyr	AAG Lys	GAC Asp	585
	TAC Tyr 180	ATC Ile	CGG Arg	GAA Glu	CGC Arg	TTC Phe 185	GAC Asp	AAT Asn	GAG Glu	ACG Thr	TTC Phe 190	CGG Arg	ATC Ile	AGC Ser	GTT Val	TAT Tyr 195	633
15	CAG Gln	GTG Val	CTC Leu	CAG Gln	GAG Glu 200	CAC His	TTG Leu	GGC Gly	AGG Arg	GAA Glu 205	TCG Ser	GAT Asp	CTC Leu	TTC Phe	CTG Leu 210	CTC Leu	681
	GAC Asp	AGC Ser	Arg	ACC Thr 215	CTC Leu	TGG Trp	GCC Ala	TCG Ser	GAG Glu 220	GAG Glu	GGC Gly	TGG Trp	CTG Leu	GTG Val 225	TTT Phe	GAC Asp	729
20	ATC Ile	ACA Thr	GCC Ala 230	Thr	AGC Ser	AAC Asn	CAC His	TGG Trp 235	GTG Val	GTC Val	AAT Asn	CCG Pro	CGG Arg 240	HIS	AAC Asn	CTG Leu	777
25	Gly	CTG Leu 245	CAG Gln	CTC Leu	TCG Ser	GTG Val	GAG Glu 250	Thr	CTG Leu	GAT Asp	GGG Gly	CAG Gln 255	Ser	ATC	AAC Asn	CCC Pro	825
	AAG Lys 260	Leu	GCG Ala	GGC Gly	CTG Leu	ATT Ile 265	Gly	CGG	CAC His	GGG Gly	CCC Pro 270	GIII	AAC	Lys	CAG Gln	Pro 275	873
30	TTC Phe	ATG Net	GTG Val	Ala	TTC Phe 280	Phe	Lys	Ala	Int	GTO	Val	. nis	THE	CGC Arg	Ser 290	ATC	921
	CGG Arg	TCC	ACC	GGG Gly 295	Ser	Lys	CAG Glm	CGC Arg	AGC Ser 300	GIN	AAC Asn	CGC Arg	TCC Ser	Lys 305		CCC Pro	969
35	AAG Lys	AAC Asn	CAC Glr 310	Gli	GCC Ala	CTG Lev	G CGC	ATC Met 315	: Ala	AAC AST	GTC Val	G GCA	GAC Glu 320	I ASI	AG0 Se1	AGC Ser	1017
40	Ser	GAC Asp 325	Gli	G AGO	G CAC	GCC Ala	TGT Cys	: Lys	AAC Lys	G CAC	GA(CTO Let 335	TA	r Va	C AGO	TTC Phe	1065

05															TAC Tyr			1113
10															TAC Tyr 370			1161
	AAC Asn	GCC Ala	ACC Thr	AAC Asn 375	CAC His	GCC Ala	ATC Ile	GTG Val	CAG Gln 380	ACG Thr	CTG Leu	GTC Val	CAC His	TTC Phe 385	ATC Ile	AAC Asn		1209
15															AAT Asn	GCC Ala		1257
															AAG Lys			1305
20				ATG H et									TAG	CTCC	TCC			1351
	GAG	AATT(CAG .	ACCC	TTTG	GG G	CCAA	GTTT:	r TC	TGGA'	TCCT	CCA:	TTGC	rcg	CCTT	GGCCAC	3	1411
	GAA	CCAG	CAG .	ACCA	ACTG	CC T	TTTG'	rgag,	A CC	TTCC	CCTC	CCT	ATCC	CCA .	ACTT	TAAAG(3	1471
25	TGT	GAGA	GTA '	TTAG	GAAA	CA T	GAGC	AGCA'	TA 1	GGCT	TTTG	ATC.	AGTT	TTT	CAGT	GGCAG	3	1531
	ATC	CAAT	GAA	CAAG.	ATCC	TA C	AAGC'	TGTG	C AG	GCAA	AACC	TAG	CAGG	AAA`.	AAAA	AACAA	3	1591
	GCA'	AAAT	GAA	TAAA	GGCC	GG G	CCAG	GTCA'	T TG	GCTG	GGAA	GTC	TCAG	CCA	TGCA	CGGAC.	r	1651
	CGT	TTCC	AGA	GGTA	ATTA	TG A	GCGC	CTAC	C AG	CCAG	GCCA	CCC	AGCC	GTG	GGAG	GAAGG	3	1711
	GGC	GTGG	CAA	GGGG	TGGG	CA C	ATTG	GTGT	C TG	TGCG	AAAG	GAA	TTAA	GAC	CCGG	AAGTT(C	1771
30	CTG	TAAT	AAA	TGTC	ACAA	TA A	AACG	AATG	A AT	GAAA	AAAA	AAA	AAAA	AAA	A			1822

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 431 amino acids
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (ix) FEATURE:

20

(D) OTHER INFORMATION: /Product="hOP1-PP"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met His Val Arg Ser Leu Arg Ala Ala Ala Pro His Ser Phe Val Ala 1 10 15

Leu Trp Ala Pro Leu Phe Leu Leu Arg Ser Ala Leu Ala Asp Phe Ser 20 25

Leu Asp Asn Glu Val His Ser Ser Phe Ile His Arg Arg Leu Arg Ser 40 45

Gln Glu Arg Arg Glu Met Gln Arg Glu Ile Leu Ser Ile Leu Gly Leu 50 60

Pro His Arg Pro Arg Pro His Leu Gln Gly Lys His Asn Ser Ala Pro

Pro His Arg Pro Arg Pro His Leu Gln Gly Lys His Asn Ser Ala Pro 65 70 75 80

Met Phe Met Leu Asp Leu Tyr Asn Ala Met Ala Val Glu Glu Gly Gly

Gly Pro Gly Gln Gly Phe Ser Tyr Pro Tyr Lys Ala Val Phe Ser

Thr Gln Gly Pro Pro Leu Ala Ser Leu Gln Asp Ser His Phe Leu Thr 115 120 125

Asp Ala Asp Met Val Met Ser Phe Val Asn Leu Val Glu His Asp Lys 130 135 140

25 Glu Phe Phe His Pro Arg Tyr His His Arg Glu Phe Arg Phe Asp Leu 145 150 150 160

Ser Lys Ile Pro Glu Gly Glu Ala Val Thr Ala Ala Glu Phe Arg Ile 165 170 175

Tyr Lys Asp Tyr Ile Arg Glu Arg Phe Asp Asn Glu Thr Phe Arg Ile 30 180 185 190

Ser Val Tyr Gln Val Leu Gln Glu His Leu Gly Arg Glu Ser Asp Leu 195 200 205

Phe Leu Leu Asp Ser Arg Thr Leu Trp Ala Ser Glu Glu Gly Trp Leu 210 215 220

Val Phe Asp Ile Thr Ala Thr Ser Asn His Trp Val Val Asn Pro Arg 225 230 235 240

His Asn Leu Gly Leu Gln Leu Ser Val Glu Thr Leu Asp Gly Gln Ser 245 250 255

Ile Asn Pro Lys Leu Ala Gly Leu Ile Gly Arg His Gly Pro Gln Asn

WO 92/07073 PCT/US91/07635

- 46 -

270 05 265 260 Lys Gln Pro Phe Met Val Ala Phe Phe Lys Ala Thr Glu Val His Phe 275 Arg Ser Ile Arg Ser Thr Gly Ser Lys Gln Arg Ser Gln Asn Arg Ser Lys Thr Pro Lys Asn Gln Glu Ala Leu Arg Met Ala Asn Val Ala Glu 10 315 Asn Ser Ser Ser Asp Gln Arg Gln Ala Cys Lys Lys His Glu Leu Tyr Val Ser Phe Arg Asp Leu Gly Trp Gln Asp Trp Ile Ile Ala Pro Glu 15 Gly Tyr Ala Ala Tyr Tyr Cys Glu Gly Glu Cys Ala Phe Pro Leu Asn Ser Tyr Met Asn Ala Thr Asn His Ala Ile Val Gln Thr Leu Val His Phe Ile Asn Pro Glu Thr Val Pro Lys Pro Cys Cys Ala Pro Thr Gln 20 390 395 Leu Asn Ala Ile Ser Val Leu Tyr Phe Asp Asp Ser Ser Asn Val Ile Leu Lys Lys Tyr Arg Asn Met Val Val Arg Ala Cys Gly Cys His 430 25 INFORMATION FOR SEQ ID NO:3: (2) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1929 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 30 (D TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA FEATURE: (ix) (A) NAME/KEY: CDS LOCATION: 103..1293 (B) (D) OTHER INFORMATION: /function= "osteogenic protein" /product= "mOP2-PP" /note= "mOP2 cDNA" (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

05	CCGA	CCAG	CT A	CCAG	TGGA	T GC	GCGC	CGGC	TGA	AAGT	CCG	AG A H	TG G et A 1	CT A la H	TG C	GT .rg	114
10	CCC Pro 5	GGG Gly	CCA Pro	CTC Leu	TGG Trp	CTA ' Leu '	TTG Leu	GGC Gly	CTT Leu	GCT Ala	CTG Leu 15	TGC Cys	GCG Ala	CTG Leu	GGA Gly	GGC Gly 20	162
	GGC Gly	CAC His	GGT Gly	CCC Pro	GGT Gly 25	CCC Pro	CCG Pro	CAC His	ACC Thr	TGT Cys 30	CCC Pro	CAG Gln	CGT Arg	CGC Arg	CTG Leu 35	GGA Gly	210
15	GCG Ala	CGC Arg	GAC Asp	CGG Arg 40	GAC Asp	ATG Het	CAG Gln	CGT Arg	GAA Glu 45	ATC Ile	CTG Leu	CCG Pro	GTG Val	CTC Leu 50	GGG Gly	CTA Leu	258
	CCG Pro	GGA Gly	CGC Arg 55	CCC Pro	GAC Asp	CCC Pro	GTG Val	CAC His 60	AAC Asn	CCG Pro	CCG Pro	CTG Leu	CCC Pro 65	GGC Gly	ACG Thr	CAG Gln	306
20	CGT Arg	GCG Ala 70	CCC Pro	CTC Leu	TTC Phe	ATG Met	TTG Leu 75	GAC Asp	CTA Leu	TAC Tyr	CAC His	GCC Ala 80	ATG Met	ACC Thr	GAT Asp	GAC Asp	354
25	Asp	GAC Asp	GGC Gly	GGG Gly	CCA Pro	CCA Pro 90	CAG Gln	GCT Ala	CAC His	TTA Leu	GGC Gly 95	wra	GCC Ala	GAC Asp	CTG Leu	GTC Val 100	402
	ATG Het	AGC Ser	TTC Phe	GTC Val	AAC Asn 105	Het	GTG Val	GAA Glu	CGC Arg	GAC Asp 110	Arg	ACC	CTG	GGC	TAC Tyr 115	CAG	450
30	GAG Glu	CCA Pro	CAC	TGG Trp 120	Lys	GAA Glu	TTC Phe	CAC His	TTT Phe 125	Asp	CTA Leu	ACC Thr	CAG Gln	ATC 1le		GCT Ala	498
	GGG Gly	GAG Glu	GCT Ala 135	Val	ACA Thr	GCT Ala	GCT Ala	GAG Glu 140	rne	CGC Arg	G ATO	TAC Tyr	Lys 145	GIL	CCC Pro	AGC Ser	546
	ACC Thi	CAC His	Pro	CTC Lev	AAC ASI	ACA Thr	ACC Thr 155	Leu	CAC His	ATO	C AGO	ATO Met	, riii	GA/	A GT(GTC Val	594
	CAA Gl: 16!	ı Glı	G CAC	TC(S Sei	AA(AGG Arg 170	g GIU	TC1	GAC Asj	TT(G TT(u Pho 17!	2 1116	r TT(e Le	G GA:	r CT	CAG Gln 180	642
	ACC Th:	G CT	C CGA	A TC	r GG(r Gl)	y Asj	GAC Glu	GG(Gl)	TG(Tr	G CT p Le	u va	G CT(G GA	C ATO	C AC e Th	A GCA r Ala 5	690
	GC	C AG	T GA	C CG.	A TG	G CT(G CT	G AA	C CA	T CA	C AA	G GA	C CT	G GG	A CT	C CGC	738

05	Ala	Ser	Asp	Arg 200	Trp	Leu	Leu	Asn	His 205	His	Lys	Asp	Leu	Gly 210	Leu	Arg	
					ACC Thr												786
10					CGA Arg												834
15					GCC Ala												882
					AGG Arg 265												930
20					CCA Pro												978
					CGC Arg												1026
25					TGG Tr											TAC r Tyr	1074
3 0					TGT Cys												1122
					TTG Leu 345												1170
3 5					TGC Cys												1218
				Asp					Val							AAC Asn	1266
			Val		GCC Ala						GGCC	CCG	CCCA	GCAT	СС		1313
	TGC	TTCT.	ACT	ACCT	TACC.	AT C	TGGC	CGGG	c cc	CTCT	CCAG	AGG	CAGA	AAC	CCTT	CTATGT	1373

05	TATCATAGCT CAGACAGGGG CAATGGGAGG CCCTTCACTT CCCCTGGCCA CTTCCTGCTA	
	AATTCTGGT CTTTCCCAGT TCCTCTGTCC TTCATGGGGT TTCGGGGCTA TCACCCCGCC	
	TCTCCATCC TCCTACCCCA AGCATAGACT GAATGCACAC AGCATCCCAG AGCTATGCTA	
•	ACTGAGAGGT CTGGGGTCAG CACTGAAGGC CCACATGAGG AAGACTGATC CTTGGCCATC	
	TCAGCCCAC AATGGCAAAT TCTGGATGGT CTAAGAAGCC CTGGAATTCT AAACTAGATG	
10	ATCTGGGCTC TCTGCACCAT TCATTGTGGC AGTTGGGACA TTTTTAGGTA TAACAGACAC	
	ATACACTTAG ATCAATGCAT CGCTGTACTC CTTGAAATCA GAGCTAGCTT GTTAGAAAAA	
	GAATCAGAGC CAGGTATAGC GGTGCATGTC ATTAATCCCA GCGCTAAAGA GACAGAGACA	
	GGAGAATCTC TGTGAGTTCA AGGCCACATA GAAAGAGCCT GTCTCGGGAG CAGGAAAAAA	
	AAAAAAACG GAATTC	
15	2) INFORMATION FOR SEQ ID NO:4:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 397 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear 	
20	(ii) MOLECULE TYPE: protein	
	<pre>(ix) FEATURE: (D) OTHER INFORMATION: /Product= "mOP2-PP"</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
25	Net Ala Met Arg Pro Gly Pro Leu Trp Leu Leu Gly Leu Ala Leu Cys 1 10 15	
	Ala Leu Gly Gly Gly His Gly Pro Gly Pro Pro His Thr Cys Pro Gln 20 25 30	
	Arg Arg Leu Gly Ala Arg Asp Arg Asp Met Gln Arg Glu Ile Leu Pro 35 40 45	
30	Val Leu Gly Leu Pro Gly Arg Pro Asp Pro Val His Asn Pro Pro Leu 50 55 60	
	Pro Gly Thr Gln Arg Ala Pro Leu Phe Het Leu Asp Leu Tyr His Ala 65 70 75 80	
	Het Thr Asp Asp Asp Gly Gly Pro Pro Gln Ala His Leu Gly Arg 85 90 95	

05	Ala	Asp	Leu	Val 100	Met	Ser	Phe	Val	Asn 105	Het	Val	Glu	Arg	Asp 110	Arg	Thr
	Leu	Gly	Tyr 115	Gln	Glu	Pro	His	Trp 120	Lys	Glu	Phe	His	Phe 125	Asp	Leu	Thr
10	Gln	Ile 130	Pro	Ala	Gly	Glu	Ala 135	Val	Thr	Ala	Ala	Glu 140	Phe	Arg	Ile	Tyr
	Lys 145	Glu	Pro	Ser	Thr	His 150	Pro	Leu	Asn	Thr	Thr 155	Leu	His	Ile	Ser	Met 160
	Phe	Glu	Val	Val	Gln 165	Glu	His	Ser	Asn	Arg 170	Glu	Ser	Asp	Leu	Phe 175	Phe
15	Leu-	Asp	Leu	Gln 180	Thr	Leu	Arg	Ser	Gly 185	Asp	Glu	Gly	Trp	Leu 190	Val	Leu
	Asp	Ile	Thr 195	Ala	Ala	Ser	Asp	Arg 200	Trp	Leu	Leu	Asn	His 205	His	Lys	Asp
20	Leu	Gly 210	Leu	Arg	Leu	Tyr	Val 215	Glu	Thr	Ala	Asp	Gly 220	His	Ser	Het	Asp
	Pro 225	Gly	Leu	Ala	Gly	Leu 230	Leu	Gly	Arg	Gln	Ala 235	Pro	Arg	Ser	Arg	Gln 240
	Pro	Phe	Met	Val	Thr 245	Phe	Phe	Arg	Ala	Ser 250	Gln	Ser	Pro	Val	Arg 255	Ala
25	Pro	Arg	Ala	Ala 260	Arg	Pro	Leu	Lys	Arg 265	Arg	Gln	Pro	Lys	Lys 270	Thr	Asn
	Glu	Leu	Pro 275		Pro	Asn	Lys	Leu 280	Pro	Gly	Ile	Phe	Asp 285	Asp	Gly	His
30	Gly	Ser 290		Gly	Arg	Glu	Val 295		Arg	Arg	His	Glu 300	Leu	Tyr	Val	Arg
	Phe 305		, Asp	Leu	Gly	Trp 310		Asp	Trp	Val	11e 315	Ala	Pro	Gln	Gly	Tyr 320
	Ser	Ala	Туг	Туг	Cys 325	Glu	Gly	Glu	Cys	330	Phe	Pro	Let	ı Asp	Ser 335	Cys
35	Het	Asn	Ala	Thr 340		h His	Ala	Ile	Let 345	ı Glr	Ser	Leu	\Val	350	Leu	Met
	Lys	Pro	Ası 35		l Val	Pro	Lys	360		s Cys	Ala	Pro	Th: 36!	r Lys	Leu	Ser
	Ala	Th: 37(r Sea	r Val	l Le	1 Ту1	Tyr 375	. Ası	Se	r Sei	r Ası	1 Ast 380	ı Va:	l Ile	e Let	Arg

05	Lys His Arg Asn Met Val Val Lys Ala Cys Gly Cys His 385 390 395	
	(2) INFORMATION FOR SEQ ID NO:5:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1941 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA	
15	(vi) ORIGINAL SOURCE: (A) ORGANISM: HOMO SAPIENS (F) TISSUE TYPE: HIPPOCAMPUS	
20	<pre>(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 5071703 (D) OTHER INFORMATION: /function= "OSTEOGENIC PROTEIN"</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:	
25	GGAATTCCGG CCACAGTGGC GCCGGCAGAG CAGGAGTGGC TGGAGGAGCT GTGGTTGGAG	60
	CAGGAGGTGG CACGGCAGGG CTGGAGGGCT CCCTATGAGT GGCGGAGACG GCCCAGGAGG	120
	CGCTGGAGCA ACAGCTCCCA CACCGCACCA AGCGGTGGCT GCAGGAGCTC GCCCATCGCC	180
	CCTGCGCTGC TCGGACCGCG GCCACAGCCG GACTGGCGGG TACGGCGGCG ACAGAGGCAT	240
	TGGCCGAGAG TCCCAGTCCG CAGAGTAGCC CCGGCCTCGA GGCGGTGGCG TCCCGGTCCT	300
30	CTCCGTCCAG GAGCCAGGAC AGGTGTCGCG CGGCGGGGCT CCAGGGACCG CGCCTGAGGC	360
50	CGGCTGCCCG CCCGTCCCGC CCCGCCCGC CGCCCGAGCC CAGCCTCCTT	420
	GCCGTCGGGG CGTCCCCAGG CCCTGGGTCG GCCGCGGAGC CGATGCGCGC CCGCTGAGCG	480
	CCCCAGCTGA GCGCCCCGG CCTGCC ATG ACC GCG CTC CCC GGC CCG CTC TGG Met Thr Ala Leu Pro Gly Pro Leu Trp 1 5	533
35	CTC CTG GGC CTG GCG CTA TGC GCG CTG GGC GGC GGC CCC GGC CTG Leu Leu Gly Leu Ala Leu Cys Ala Leu Gly Gly Gly Pro Gly Leu 10 25	581
	CGA CCC CCG CCC GGC TGT CCC CAG CGA CGT CTG GGC GCG CGC GAC CGG	629

05	Arg	Pro	PTO	PTO	30	cys	Pro	GIN	Arg	35	Leu	61À	Ala	Arg	Asp 40	Arg	
					GAG Glu												677
10					CCÀ Pro												725
15					GAC Asp												773
					GAG Glu												821
20					GAG Glu 110												869
					CGC Arg												917
2 5					GAG Glu												965
30					CTC Leu												1013
					TCT Ser												1061
35					GGC Gly 190												1109
			Leu		AAG Lys												1157
40					GGG Gly												1205
					CCA Pro												1253

05		235					240					245					
	AGG Arg 250	GCC Ala	AGT Ser	CCG Pro	AGT Ser	CCC Pro 255	ATC Ile	CGC Arg	ACC Thr	CCT Pro	CGG Arg 260	GCA Ala	GTG Val	AGG Arg	CCA Pro	CTG Leu 265	1301
10	AGG Arg	AGG Arg	AGG Arg	CAG Gln	CCG Pro 270	AAG Lys	AAA Lys	AGC Ser	AAC Asn	GAG Glu 275	CTG Leu	CCG Pro	CAG Gln	GCC Ala	AAC Asn 280	CGA Arg	1349
	CTC Leu	CCA Pro	GGG Gly	ATC Ile 285	TTT Phe	GAT Asp	GAC Asp	GTC Val	CAC His 290	GGC Gly	TCC Ser	CAC His	GGC Gly	CGG Arg 295	CAG Gln	GTC Val	1397
15	TGC Cys	CGT Arg	CGG Arg 300	CAC His	GAG Glu	CTC Leu	TAC Tyr	GTC Val 305	AGC Ser	TTC Phe	CAG Gln	GAC Asp	CTC Leu 310	GGC Gly	TGG Trp	CTG Leu	1445
2 0	GAC Asp	TGG Trp 315	GTC Val	ATC Ile	GCT Ala	CCC Pro	CAA Gln 320	GGC Gly	TAC Tyr	TCG Ser	GCC Ala	TAT Tyr 325	TAC Tyr	TGT Cys	GAG Glu	GGG Gly	1493
	GAG Glu 330	TGC Cys	TCC Ser	TTC Phe	CCA Pro	CTG Leu 335	GAC Asp	TCC Ser	TGC Cys	ATG Met	AAT Asn 340	Ala	ACC Thr	AAC Asn	CAC His	GCC Ala 345	1541
25	ATC Ile	CTG Leu	CAG Gln	TCC Ser	CTG Leu 350	GTG Val	CAC His	CTG Leu	ATG Met	AAG Lys 355	CCA Pro	AAC Asn	GCA Ala	GTC Val	CCC Pro 360	AAG Lys	1589
	GCG Ala	TGC Cys	Cys	GCA Ala 365	CCC Pro	ACC Thr	AAG Lys	CTG Leu	AGC Ser 370	GCC Ala	ACC Thr	TCT Ser	GTG Val	CTC Leu 375	TAC Tyr	TAT Tyr	1637
30	GAC Asp	AGC Ser	AGC Ser 380	Asn	AAC Asn	GTC Val	ATC Ile	CTG Leu 385	Arg	AAA Lys	GCC Ala	CGC Arg	AAC Asn 390	Het	GTG Val	GTC Val	1685
35				GGC Gly			TGA	GTCA	GCC	CGCC	CAGC	CC T	ACTG	CAGC	A		1733
	TTA	CACT	GGC	CGTC	GTTT	TA C	AACG'	TGTG	A CT	GGGA	AAAC	CCT	GGCG	ATT	CCCA	ACTTAA	1793
	TCG	CCTT	GCA	GCAC	ATCC	cc c	TTTC	GCCA	G CT	GGCT	ATAA	GCG	AAGA	GGC	CCCG	CACCGA	1853
	TCG	CCCT	TCC	CAAC	AGTT	GC G	CCCC	AGTG	A AT	GGCG	AATG	GCA	TTAA	GTA	AGCG	ATAATA	1913
	TTT	TGTT	AAA	ATTC	GCGT	TA A	TTTA	TTT									1941

05			(i)	SE(A (B (D) T	ENGT:	H: 3 ami	CTER 99 au no a lin	mino cid	CS: aci	ds					
			(ii)	но	LECU	LE T	YPE:	pro	tein							
10		(ix)	FE (D	ATUR	E: Ther	INF	ORMA	TION	: /p	rodu	ct=	"hOP	2 -PP	Ħ	
			(xi)	SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:6:				
	Met 1	Thr	Ala	Leu	Pro 5	Gly	Pro	Leu	Trp	Leu 10	Leu	Gly	Leu	Ala	Leu 15	Cys
15	Ala	Leu	Gly	Gly 20	Gly	Gly	Pro	Gly	Leu 25	Arg	Pro	Pro	Pro	Gly 30	Cys	Pro
	Gln	Arg	Arg 35	Leu	Gly	Ala	Arg	Asp 40	Arg	Asp	Val	Gln	Arg 45	Glu	Ile	Leu
20	Ala	Val 50	Leu	Gly	Leu	Pro	Gly 55	Arg	Pro	Arg	Pro	Arg 60	Ala	Pro	Pro	Ala
	Ala 65	Ser	Arg	Leu	Pro	Ala 70	Ser	Ala	Pro	Leu	Phe 75	Het	Leu	Asp	Leu	Tyr 80
	His	Arg	Met	Ala	Gly 85	Asp	Asp	Asp	Glu	Asp 90	Gly	Ala	Ala	Glu	Ala 95	Leu
2 5	Gly	Arg	Ala	Asp 100	Leu	Val	Het	Ser	Phe 105	Val	Asn	Met	Val	Glu 110	Arg	Asp
	Arg	Ala	Leu 115	Gly	His	Gln	Glu	Pro 120	His	Trp	Lys	Glu	Phe 125	Arg	Phe	Asp
30	Leu	Thr	Gln	Ile	Pro	Ala	Gly 135	Glu	Ala	Val	Thr	Ala 140	Ala	Glu	Phe	Arg
	Ile 145		Lys	Val	Pro	Ser 150	Ile	His	Leu	Leu	Asn 155	Arg	Thr	Leu	His	Val 160
	Ser	Het	: Phe	Gln	Val 165	Val	Gln	Glu	Gln	Ser 170	Asn	Arg	Glu	Ser	Asp 175	Leu
35	Phe	Phe	e Leu	Asp 180	Leu	Gln	Thr	Leu	Arg 185	Ala	Gly	Asp	Glu	Gly 190	Trp	Leu
	Va]	Let	1 Asp 195		Thr	Ala	Ala	Ser 200	Asp	Cys	Trp	Leu	Leu 205	Lys	Arg	, His
	Lys	5 Asj 210	p Le v	ı Gly	/ Leu	ı Arg	Let 21:	ı Tyı	· Val	Glu	ı Thi	Glu 220	Asp	Gly	His	S Set

05	Val 225	Asp	Pro	Gly	Leu	Ala 230	Gly	Leu	Leu	Gly	Gln 235	Arg	Ala	Pro	Arg	Ser 240
	Gln	Gln	Pro	Phe	Val	Val 245	Thr	Phe	Phe	Arg 250	Ala	Ser	Pro	Ser	Pro 255	Ile
10	Arg	Thr		Arg 260	Ala	Val	Arg	Pro	Leu 265	Arg	Arg	Arg	Gln	Pro 270	Lys	Lys
	Ser	Asn	Glu 275	Leu	Pro	Gln	Ala	Asn 280	Arg	Leu	Pro	Gly	Ile 285	Phe	Asp	Asp
	Val	His 290	Gly	Ser	His	Gly	Arg 295	Gln	Val	Cys	Arg	Arg 300	His	Glu	Leu	Tyr
15	Val 305	Ser	Phe	Gln	Asp	Leu 310	Gly	Trp	Leu	Asp	Trp 315	Val	Ile	Ala	Pro	Gln 320
	Gly	Tyr	Ser	Ala	Tyr 325	Tyr	Cys	Glu	Gly	Glu 330	Cys	Ser	Phe	Pro	Leu 335	Asp
20	Ser	Cys		Asn 340	Ala	Thr	Asn	His	Ala 345	Ile	Leu	Gln	Ser	Leu 350	Val	His
	Leu	Ket	Lys 355	Pro	Asn	Ala	Val	Pro 360	Lys	Ala	Cys	Cys	Ala 365	Pro	Thr	Lys
	Leu	Ser 370	Ala	Thr	Ser	Val	Leu 375	Tyr	Tyr	Asp	Ser	Ser 380	Asn	Asn	Val	Ile
2 5	Leu 385	Arg	Lys	Ala	Arg	Asn 390	Met	Val	Val	Lys	Ala 395	Cys	Gly	Cys	His	
	121	TNI	FORM	ለ ጥፕ ሰነ	J FOI	S SEC	חז ר	NO.	7 •							

- - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 102 amino acids
 - TYPE: amino acid (B)
 - TOPOLOGY: linear (D)
 - (ii) MOLECULE TYPE: protein
 - (ix) FEATURE:

30

- (A) NAME/KEY: Protein (B) LOCATION: 1..102
- (D) OTHER INFORMATION: /label= OPX /note= "WHEREIN EACH XAA IS INDEPENDENTLY SELECTED FROM A GROUP OF ONE OR MORE SPECIFIED AMINO ACIDS AS DEFINED IN THE SPECIFICATION
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

	С	ys 1	Xaa	Xaa	His	Glu 5	Leu	Tyr	Val	Xaa	Phe 10	Xaa	Asp	Leu	Gly	Trp 15	Xaa	
5	A	sp	Trp	Xaa	Ile 20	Ala	Pro	Xaa	Gly	Tyr 25	Xaa	Ala	Tyr	Tyr	Cys 30	Glu	Gly	
	G	lu	Cys	Xaa 35	Phe	Pro	Leu	Xaa	Ser 40	Xaa	Het	Asn	Ala	Thr 45	Asn	His	Ala	
10	1	le	Xaa 50	Gln	Xaa	Leu	Val	His 55	Xaa	Xaa	Xaa	Pro	Xaa 60	Xaa	Val	Pro	Lys	ı
		laa 55	Cys	Cys	Ala	Pro	Thr 70	Xaa	Leu	Xaa	Ala	Xaa 75	Ser	Val	Leu	Tyr	Xa a 80))
15	I	Asp	Xaa	Ser	Xaa	Asn 85	Val	Xaa	Leu	Xaa	Lys 90	Xaa	Arg	Asn	Het	Val 95	. Va]	Ĺ
20	3	Kaa	Ala	Cys	Gly 100		His											
	(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:8	:									
25			(i)	SE (A (B) I	ENGT YPE:	H: 9 ami	CTER 7 am no a lin	ino cid	CS: acid	ls							
30			(ii)	MC	LECU	LE I	YPE:	pro	tein	1								
35			(ix)	(<i>I</i> (I	3í I	IAME/	ION: R INI /not ONE	Prof. 1 FORMA TORMA TORMA	97 TION WHEI	N: /] REIN 20 N/	EACI ATUR	H XAA ALLY-	A INI -OCCI	JEPE JRRI	NG L	-120	nek,	CATES
40			(xi) Si	EQUE	NCE I	DESC	RIPT:	ION:	SEQ	ID 1	NO:8	:					
		3	Kaa i	Xaa :	Xaa 🛚	Xaa :	Xaa : 5	Xaa :	Xaa :	Xaa :	Xaa :	Xaa : 10	Xaa :	Xaa	Xaa	Xaa	Xaa 15	Xaa
45		3	Kaa	Xaa	Xaa :	Xaa : 20	Xaa :	Xaa 🗆	Xaa i	Xaa	Cys 25	Xaa :	Xaa	Xaa	Cys	Xaa 30	Xaa	Xaa
50		3	Xaa	Xaa	Xaa 35	Cys	Xaa	Xaa	Xaa	Xaa 40	Xaa	Xaa	Xaa	Xaa	Xaa 45	Xaa	Xaa	Xaa
		:	Xaa	Xaa 50	Xaa	Xaa	Xaa	Xaa	Xaa 55	Xaa	Xaa	Xaa	Xaa	Xaa 60	Cys	Cys	Xaa	Xaa

05 Xaa INFORMATION FOR SEQ ID NO:9: 10 (2) (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 102 amino acids (B) TYPE: amino acid TOPOLOGY: linear (D) (ii) MOLECULE TYPE: protein 15 FEATURE: (ix) (A) NAME/KEY: Protein LOCATION: 1..102 **(B)** (D) OTHER INFORMATION: /label= PROTEIN /note= "WHEREIN EACH XAA INDEPENDENTLY INDICATES 20 ONE OF THE 20 NATURALLY-OCCURRING L-ISOMER A-AMINO AICDS, OR A DERIVATIVE THEREOF." (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9: 25 Xaa Cys Xaa Xaa Xaa Xaa Xaa Cys Xaa Xaa Xaa Xaa Xaa Xaa Xaa 30 Xaa Xaa Cys Xaa Cys Xaa

05 What is claimed is:

- 1. A polypeptide chain comprising an amino acid sequence described by residues 303-399 of Seq. ID No. 5.
- The polypeptide chain of claim 1 comprising an
 amino acid sequence described by residues 297-399 of
 Seq. ID No. 5.
 - 3. The polypeptide chain of claim 2 comprising of amino acid sequence described by residues 267-399 of Seq. ID No. 5.
- 4. The polypeptide chain of claim 3 comprising an amino acid sequence described by residues 264-399 of Seq. ID No. 5.
- The polypeptide chain of claim 4 comprising an amino acid sequence described by residues 240-399 of
 Seq. ID No. 5.
 - 6. The polypeptide chain of claim 5 comprising an amino acid sequence described by residues 1-399 of Seq. ID No. 5.
- 7. A polypeptide chain comprising an amino acid sequence described by residues of 301-397 of Seq. ID No. 3.
 - 8. The polypeptide chain of claim 7 comprising an amino acid sequence described by residues 296-397 of Seq. ID No.3.

- 9. The polypeptide chain of claim 8 comprising an amino acid sequence described by residues 259-397 of Seq. ID No. 3.
- 10. The polypeptide chain of claim 9 comprising an amino acid described by residues 1-397 of Seq. ID

 No. 3.
- 11. A polypeptide chain useful as a subunit of a dimeric osteogenic protein comprising a pair of disulfide-bonded polypeptide chains, said polypeptide chain having an amino acid sequence described by residues 303-399 of Seq. ID No. 5, including allelic and species variants thereof, such that the dimeric osteogenic protein comprising said polypeptide chain has a conformation capable of inducing endochondral bone formation when implanted in a mammal in association with a matrix.
 - 12. The polypeptide chain of claim 11 wherein said amino acid sequence comprises residues 261-399 of Seq. ID 5.
- 13. The polypeptide chain of claim 11 wherein the
 amino acid sequence comprises residues 301-397 of Seq.
 ID No. 3.
 - 14. The polypeptide chain of claim 13 wherein said amino acid sequent comprises residues 259-397 of Seq. ID No. 3.
- 30 15. A dimeric osteogenic protein capable of inducing endochondral bone formation in a mammal when implanted in said mammal in association with a matrix;

WO 92/07073

- os said protein comprising a pair of disulfide-bonded polypeptide chains constituting a dimeric species, wherein each said polypeptide chain is the polypeptide chain of claim 11.
- 16. The polypeptide chain of claim 3 or 11 produced by expression of recombinant DNA in a host cell.
 - 17. The polypeptide chain of claim 16 wherein said host cell is a procaryotic host cell.
 - 18. The polypeptide chain of claim 16 wherein said host cell is a mammalian cell.
- 19. The polypeptide of claim 1, 3 or 11 that is glycosylated.
 - 20. A nucleic acid encoding the polypeptide chain of claim 1, 3, or 11.
- 21. A dimeric protein comprising a pair of polypeptide chains expressed from a DNA sequence described by ID No. 3 or ID No. 5, including allelic and species variants thereof, such that, when said polypeptide chains are oxidized to produce a disulfide-bonded dimeric species, the dimeric species has a conformation that is capable of inducing endochondral bone or cartilage formation when disposed within a matrix and implanted in a mammal.

hOP2 mOP2	Ala	Val Ala	Arg	Pro	Leu 5	Arg Lys	Arg	Arg
hOP2 mOP2	Gln	Pro 10	Lys	Lys	Ser Thr	Asn	Glu 15	Leu
hOP2 mOP2	Pro	Gln His	Ala Pro	Asn 20	Arg Lys	Leu	Pro	Gly
hOP2 mOP2	Ile '25	Phe	Asp	Asp 	Val Gly	His 30	Gly	Ser
hOP2 mOP2	His Arg	Gly	Arg 35	Gln Glu	Val	Cys	Arg	Arg 40
hOP2 mOP2	His	Glu	Leu	Tyr	Val 45	Ser Arg	Phe	Gln Arg
hOP2 mOP2	Asp	Leu 50	Gly	Trp	Leu	Asp	Trp 55	Val
hOP2 mOP2	Ile	Ala	Pro	Gin 60	Gly	Tyr	Ser	Ala
hOP2 mOP2	Tyr 65	Tyr	Cys	Glu	Gly	Glu 70	Cys	Ser Ala

Fig. 1.1

hOP2 mOP2	Phe	Pro	Leu 75	Asp	Ser	Cys	Met	Asn 80
			75					
hOP2 mOP2	Ala	Thr	Asn	His		Ile	Leu	Gin
					85			
hOP2 mOP2								
HOL Z	• • •	90	• • •	• • •	• • •	• • •	95	•••
hOP2	Asn	Ala	Val	Pro	Lys	Ala	Cys	Cys
mOP2	Asp	Val	• • •	100	• • •	• • •	• • •	• • •
hOP2 mOP2			Thr	Lys	Leu	Ser	Ala	Thr
MOP 2	105	• • •	• • •	•••	• • •	iio	• • •	• • •
hOP2	Ser	Val	Leu	Tyr	Tyr	Asp	Ser	Ser
mOP2	• • •	• • •	115	•••	• • •	• • •	• • •	120
hOP2	Asn	Asn	Val	Ile	Leu	Arg	Lys	Ala
mOP2	•••	• • •	• • •	• • •	125	• • •	• • •	птр
hOP2	Arg		Met	Val	Val	Lys	Ala	Cys
mOP2	• • •	130	• • •	• • •	• • •	• • •	135	• • •
hOP2 mOP2	Gly	Cys	Hi	S				
	• • •		• •	•				

Fig. 1.2

SUBSTITUTE SHEET

hOP1 mOP1 hOP2 mOP2	 Ala	 Val	Arg	Gly Pro	Leu	Gln Arg Lys	• • •	 Arg	• • •
hOP1 mOP1 hOP2 mOP2	Pro	Lys	Lys	Ser	 Asn	Pro Glu Glu 15	 Leu	Pro	Gln
hOP1 mOP1 hOP2 mOP2	 Ala	 Asn	 Arq	Leu	Pro	Ala Gly Gly	Ser Ile	 Phe	 Asp
hOP1 mOP1 hOP2 mOP2	Asp	 Val	His	Gly	• • •	Asp His Arg	Gly	• • •	• • •
hOP1 mOP1 hOP2 mOP2	 Val	• • •	 Arg	 Arg	• • •	Glu	• • •	• • •	• • •

Fig. 2.1

hOP1 mOP1 hOP2 mOP2	• • •		Gln	• • •	• • •	•••	• • •	Leu	• • •
hOP1 mOP1 hOP2 mOP2	• • •	Ile Val Val	• • •	• • •	• • •	Gln	• • •	• • •	Ser
hOP1 mOP1 hOP2 mOP2	Ala	• • •	•••	_	• • •	• • •		• • •	Ser
hOP1 mOP1 hOP2 mOP2	• • •	Pro	• • •	 Asp	• • •	Cys	• • •	• • •	• • •
hOP1 mOP1 hOP2 mOP2	Thr	Asn	His	• • •	• • •	 Leu		Ser	• • •

Fig. 2.2

hOP1 mOP1 hOP2 mOP2	• • •	• • •	 Leu	 Met	Lys	• • •	Glu Asp Asn Asp	 Ala	• • •
hOP1 mOP1 hOP2 mOP2	• • •	• • •	 Ala	• • •	• • •	• • •	Pro	• • •	 Lys
hOP1 mOP1 hOP2 mOP2	• • •	Ser	• • •	Thr	• • •	• • •	Leu 115	• • •	Tyr
hOP1 mOP1 hOP2 mOP2	• • •	Ser	• • •	 Asn	• • •	• • •	Ile Asp 	• • •	Arg
hOP1 mOP1 hOP2 mOP2	•••	•••	• • •	• • •	• • •	• • •	Val 	Lys	
hOP1 mOP1 hOP2 mOP2	Ala 135	Cys	Gly	Cys	His				

Fig. 2.3

INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 91/07635

	ECT MATTER (if several classification sym		
	t Classification (IPC) or to both National Clas C. 12 N 15/00 C 07	sification and IPC K 7/10 C 07 K 13/	'n
Int.C1.5 A 61 K 37/02		K //10 C C K 13/	· · · · · · · · · · · · · · · · · · ·
II. FIELDS SEARCHED			
	Minimum Document		
Classification System	CI	assification Symbols	
Int.Cl.5	C 07 K A	61 K	
	Documentation Searched other the to the Extent that such Documents are	an Minimum Documentation e Included in the Fields Searched ⁶	
	ED TO DE DELEVIA T		
III. DOCUMENTS CONSIDER		e of the volument passages 12	Relevant to Claim No.13
Category ° Citation of I	Occument, 11 with indication, where appropriate	e, or the relevant passages	Activities to Class 110.
X WO,A,8 19 Oc	8909788 (CREATIVE BIOMOL tober 1989, see the whole	LECULES) e dcument	11,15- 21 1-10,12 -14
X WO,A,	9011366 (GENETICS INSTIT er 1990, see the whole do	TUTE) 4 ocument	11,15- 21
A			11,15- 21
Unive Ozkay prote	Journal, volume 9, no. 7, rsity Press (Eynsham, Oxinak et al.: "OP-1 cDNA er in in the TGF-beta familia 2093, see the whole artic	ford, GB) E. ncodes an osteogenic iy", pages	11,15 - 21
	10		vices filing date
considered to be of part "E" earlier document but pu filling date "L" document which may th which is cited to establis citation or other special "O" document referring to a other means	eneral state of the art which is not icular relevance blished on or after the international row doubts on priority claim(s) or is the publication date of another reason (as specified) n oral disclosure, use, exhibition or	"T" later document published after the interna or priority date and not in conflict with th cited to understand the principle or theory invention "X" document of particular relevance; the clai cannot be considered novel or cannot be of involve an inventive step "Y" document of particular relevance; the clai cannot be considered to involve an invent document is combined with one or more of ments, such combination being obvious to in the art.	ne application but y underlying the med invention considered to med invention ive step when the wher such docu-
"P" document published price later than the priority d	or to the international filing date but ate claimed	"&" document member of the same patent fam	nily
IV. CERTIFICATION			ah Danari
Date of the Actual Completion of		Date of Mailing of this International Seat	B 1992
International Searching Authoric	y EAN PATENT OFFICE	Signature of Authorized Offices	F. TAZELAAK

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9107635 SA 53017

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 12/02/92

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent document cited in search report	Publication date	Pater men	Publication date	
WO-A- 8909788	19-10-89	US-A- US-A- AU-A- EP-A- EP-A- JP-T- WO-A- AU-A- EP-A- JP-T- WO-A- US-A-	4968590 5011691 3444989 3530589 0372031 0362367 3500655 3502579 8909787 5174790 0411105 3504736 9010018 4975526	06-11-90 30-04-91 03-11-89 03-11-89 13-06-90 11-04-90 14-02-91 13-06-91 19-10-89 26-09-90 06-02-91 17-10-91 07-09-90 04-12-90
WO-A- 9011366	04-10-90	AU-A- CA-A- EP-A-	5357790 2030518 0429570	22-10-90 29-09-90 05-06-91